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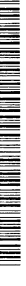
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METHOD OF IDENTIFYING AND QUANTIFYING SPECIFIC FUNGI AND BACTERIA

Cross-reference to Related Applications

The present application is a continuation in part of Serial No. 09/290,990, filed April 14, 1999, which claims priority from provisional application Serial No. 60/081,773, filed April 15, 1998, the entire contents of both of which are hereby incorporated by reference.

Field of the Invention

The present invention relates to a method of identifying and quantifying specific fungi and bacteria using specific DNA sequences, as described and taught herein. These sequences can be used with real time detection of PCR products with a fluorogenic probe system or other molecular probes like hybridization.

Background of the Invention

Fungi and bacteria are the source of or contribute to many health problems including infections, gastroenteritis, ulcers, asthma, allergies and sinusitis. The rapid identification of the microorganisms is critical for diagnosis and treatment. In addition, detecting and/or quantifying these microorganisms in the environment may help to prevent adverse health effects.

Limitations of Current Technology

In order to determine the risk fungi and bacteria pass to human health it is necessary to know when fungi and bacteria are present and in what numbers. Fungi and bacteria can be ingested, inhaled, or might enter the body through

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abrasions or punctures. It is important to identify these microorganisms as specifically and as rapidly as possible. Some species of a particular genus are harmless whereas others of the same genus may cause significant health effects. So without knowing precisely what microorganisms are present and in what numbers, it is impossible to evaluate the potential for negative health effects or the establishment of a risk assessment.

In the past, the detection and quantitative measurement of fungi and bacteria in samples has been performed either by direct microscopic examination of the collected cells or by growing cells on a suitable medium and identification and enumeration of the resultant colonies. The first method is highly labor intensive and is subject to potential errors in the recognition and positive identification. The second method is both time consuming and subject to significant quantitative inaccuracy. Both methods require extensive experience on the part of the analyst.

Some molecular approaches, such as the conventional polymerase chain reaction (PCR) procedure, are subject to inaccuracies due to the difficulty of quantifying the product. This procedure is also relatively slow and requires expertise in molecular biology.

Summary of the Invention

It is an object of the present invention to provide a simple, reliable method for detecting and quantifying some fungs and bacteria by using the nucleotide sequences specific to each species or group of species of funci and bacteria, as described herein.

According to the present invention, fungi and bacteria can be identified and quantified by using a nucleotide sequence specific to the particular species or, in the case of some fungi, group of species. Many methods including using real time, probe-based detection of polymerase chain reaction (PCR) products (e.g. TaqManTM system) or other methods of detection and quantification including hybridization or conventional PCR could be used with these sequences.

THEORY

Each microorganism is unique because of the sequence of some of the nucleatides in its DNA. However, there are many sequences which are common to more than one organism. There is thus a hierarchy or classification into which all microorganisms can be arranged. The "species" are typically the finest level of distinction that is recognized for separation of different members of a given genus. In the past, species were separated on the basis of morphological or biochemical differences. In order to identify or separate different spaces on the result of the basis of morphological or sequences that are unique to a given species but at the same

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time common to all isolates of a given species.

For this invention the internal transcribed spacer (ITS) regions of nuclear ribosomal DNA (rDNA) of the different fungi were used. For the bacteria, the sequences of unique enzymes were chosen.

To apply this invention, a number of possible detection methods are possible. For example, the TaqManTM, 3'-5' exonuclease assay signals the formation of PCR amplicons by a process involving the nucleolytic degradation of a double-labeled fluorogenic probe that hybridizes to the target template at a site between the two primer recognition sequences (cf. U.S. Patent 5,876,930). The model 7700 automates the detection and quantitative measurement of these signals, which are stoichiometrically related to the quantities of amplicons produced, during each cycle of amplification. In addition to providing substantial reductions in the time and labor requirements for PCR analyses, this technology permits simplified and potentially highly accurate quantification of target sequences in the reactions.

There are additional systems and other molecular approaches that operate upon essentially the same principal. What is common to all of these technologies is the need for the identification of specific sequences that are unique to the targeted organism but common to all members of the

of the sequences in the identification and quantification of specific fungi and bacteria.

Brief Description of the Drawings

Figure 1 illustrates the sensitivity of the assay of the present invention.

Figure 2 shows the actual vs. the expected amounts of conidia detected.

Figure 3 shows TaqMan Threshold Responses from tenfold dilutions of a single DNA extract.

Figure 4 shows H. pylori counts per assay plotted against cycle threshold values.

Detailed Description of the Invention

DNA extraction

Genomic DNA is extracted using standard methods, e.g., the glass bead milling and glass milk adsorption method or any similar procedure of extracting genomic DNA.

Reactions are prepared in 0.5 ml thin-walled, optical grade PCR tubes (PE Applied Biosystems, Foster City CA) by addition of the following components: 12.5 ul of TagMan Universal Master Mix (a 2X-concentrated, proprietary mixture of AmpliTaq GoldTM DNA polymerase, AmpErase& UNG, dNTPs with UTP, passive reference dye and optimized buffer components, PE Applied Biosystems, Foster City CA); 2.5 µl of a mixture of forward and reverse primers (10 nM each); 2.5 µl of 400 nM lagMan grope; 1.1 µl of 1 mg,ml povine serum albumin, fraction V (Sigma Chemical, St. Louis, MO) and 5 µl of DNA template.

For each targeted fungus or bacterium, the appropriate forward primer, reverse primer and probe (Table 1) are to be obtained. The probe is labeled with an appropriate set of dyes or other markers for the particular system of measurement being used.

For each target species or group of species, a calibrator sample with a known number of conidia is used as a standard. To ensure that the sample matrix does not affect the FCR reaction and, thus the quantitative results, an internal standard is used. Addition of these conidia or cells to both the test and calibrator samples normalize the target species or group for potential sample to sample variability in DNA extraction efficiencies.

Table 1. List of Fungal Primers and Probes

Absidia coerulea/glauca

Forward Primer NS92F: 5'-CACCGCCCGTCGCTAC (SEQ ID NO:1)

Reverse Primer AcoerR1: 5'-TCTAGTTTGCCATAGTTCTCTTCCAG (SEQ ID NO:2)

Frode MucP1: 5'-00GATTGARTGGTTATAGTGAGCATATGGGATC (SEQ ID NO:3)

Absidia corymbifera

Forward Frimer NS90F: 5'-CACCGCCCGTCGCTAC (SEQ ID NO:4)

Acremonium strictum

Forward Primer AstroF1: 5'-CAACCCATTGTGAACTTACCAAAC (SEQ

ID NO:7)

Reverse Primer AstroR1: 5'-GGCCCCTCAGAGAAATACGATT (SEQ ID

110:8)

Probe AstroP1: 5'-TCAGCGCGCGGTGGCCTC (SEQ ID

NO:9)

Alternaria alternata

Forward Frimer AaltrF1: 5'-GGCGGGCTGGAACCTC (SEQ ID NO:10)

Reverse Frimer AltrR1-1: 5'-GCAAFTACAAAAGGTTTATGTTTGTCGTA (SEQ ID NO:11)

Reverse Frimer AaltrR1-1: 5'-TGCAATTACTAAAGGTTTATGTTGTCGTA (SEQ II NO:12)

Probe AaltrP1: 5'-TTACAGCCTTGCTGAATTATTCACCCTTGTCTTT (SEQ ID NO:13)

Apophysomyces elegans and Saksenea vasiformis

Forward Frimer NS92F: 5'-CACCGCCCGTCGCTAC (SEQ ID NO:14)

Reverse Frimer AelegR1: 5'-GACTCGAATGAGTTCTCGCTTC (SEQ ID

NC:15)

Probe AelogF1: 5'-TGGCCAAGACCAGAATATGGGATTGC

(SEQ ID NO:16)

Aspergillus flavus/oryzae

Forward Frimer AflavF1: 5'-CGAGTGTAGGGTTCCTAGCGA (SEQ ID NO:17)

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Probe AflavP1: 5'-TCCCACCCGTGTTTACTGTACCTTAGTTGCT

(SEQ ID NO:19)

Aspergillus fumigatus, Neosartorya fischeri

Forward Primer AfumiF1: 5'-GCCCGCCGTTTCGAC (SEQ ID NO:20)

Reverse Primer AfumiR1: 5'-CCGTTGTTGAAAGTTTTAACTGATTAC

(SEQ ID NO:21)

Probe AfumiP1: 5'~CCCGCCGAAGACCCCAACATG (SEQ ID

NO:22)

Aspergillus niger/foetidus/phoenicus

Forward Primer AnigrF1: 5'-GCCGGAGACCCCAACAC-3' (SEQ ID NO:23)

Reverse Primer AnigrR1: 5'-TGTTGAAAGTTTTAACTGATTGCATT-3' (SEQ ID NO:24)

Probe AnigrP1: 5'-AATCAACTCAGACTGCACGCTTTCAGACAG (SEQ ID NO:25)

Aspergillus nomius

Forward Primer AflavF1: 5'-CGAGTGTAGGGTTCCTAGCGA-3' (SEQ ID

N0:26)

Reverse Primer AnomiR1: 5'-CCGGCGGCCTTGC-3' (SEQ ID NO:27)

Probe AflavP1: 5'-TCCCACCCGTGTTTACTGTACCTTAGTTGCT

(SEQ ID NO:28)

Aspergillus ochraceus/ostianus/auricomus

Forward Frimer RichrF1: 51-ARRITECCARTOGGTGTATACC-31 -SEQ

II NO:29

Reverse Primer AcchrR1: 5'-00GG0GAG0GCTGTG-3' (SEQ ID

N0:30)

Probe AcchrP1: 5'-ACCTTGTTGCTTCGGCGAGCCC (SEQ ID

NG:31)

Reverse Primer AparaR3: 5'-GCCCGGGGGCTGACG (SEQ ID NO:33)

Probe AflavP1: 5'-TCCCACCCGTGTTTACTGTACCTTAGTTGCT

(SEQ ID No:34)

Aspergillus restrictus/caesillus/conicus

Forward Primer ArestF2: 5'-0GGGCCCTTCAT-3' (SEC ID

110:35)

Reverse Primer ArestR1: 5'-GTTGTTGAAAGTTTTTAACGATTTTTCT

(SEQ ID NO:36)

Probe ArestP1: 5'-000GCGGGAGACTCCAACATTG (SEO ID

NO:37)

Aspergillus sydowii

Forward Primer AsydoFl: 5'-CAACCTCCCACCCGTGAA (SEQ ID

110:38)

Reverse Primer versR1: 5'-CCATTGTTGAAAGTTTTGACTGATTTTA

(SEQ ID NO:39)

Probe VersF1: 5'-AGACTGCATCACTCTCAGGCATGAAGTTCAG

(SEQ ID No:40)

Aspergillus tamarii

Forward Primer AflavF1: 5'-CGAGTSTAGGGTTCCTAGCGA (SEQ ID

NO:41)

Reverse Primer AtamaR1: 5'-COCGGCGGCCTTAA (SEQ ID NO:42)

Probe AflavP1: 5'-TOGCACCOGTGTTACTGTACCTTAGTTGCT

(SEQ ID NO:43.

Aspergillus terreus

Forward Primer AterrF1: 5'-TTACCGAGTGCGGGTCTTTA (SEQ ID

NC:44)

Reverse Primer - AtomaRi E'HragasaanakaaRer seg oo Ma:451

Probe AterrP1: 5'-AACCTCCCACCCGTGACTATTGTACCTTG

(SEQ ID NO:46)

Aspergillus ustus

Forward Primer AustsF1: 5'-GATCATTACCGAGTGCAGGTCT (SEQ ID

NO:47)

Reverse Primer AustsR1: 5'-GCCGAAGCAACGTTGGTC (SEO ID

NO:48)

Probe AustsP1: 5'-CCCCCGGGCAGGCCTAACC (SEQ ID

NO:49)

Aspergillus versicolor

Reverse Primer versE1: 5'-CCATTGTTGAAAGTTTTTGACTGATTTTA

(SEQ ID NO:51)

Probe versF1: 5'-AGACTGCATCACTCTCAGGCATGAAGTTCAG

(SEQ ID NO:52)

Chaetomium globosum

Forward Primer CglobF1: 5'-CCGCAGGCCCTGAAAAG (SEO ID

NO:53)

Probe CqlobP1: 5'-AGATGTATGCTACGCTCGGTGCGACAG

(SEQ ID NO:55)

Cladosporium cladosporicides

Type 1

Forward Primer ColadIF1: 5'-CATTACMAGTGACCCCGGTCTAAC (SEQ

ID No:56)

Type 2

Forward Primer Colad2F1: 5'-TACAAGTGACCCCGGCTACG (SEQ ID

No:59)

NO:60)

Probe DoladF1: 5'-DOGGGATGTTCATAACCCTTTGTTGTCC

SEQ ID NO:61)

Cladosporium herbarum

Forward Primer CherbF1: 5'-AAGAACGCCCGGGCTT (SEQ ID NO:62)

Reverse Primer CherbR1: 5'-CGCAAGAGTTTGAAGTGTCCAC (SEQ ID

210:631

Probe CherbP1: 5'-CTGGTTATTCATAACCCTTTGTTGTCCGACTCTG

(SEQ ID NO:64)

Cladosporium sphaerospermum

NO:65)

Reverse Primer CsphaR1: 5'-GGGGTTGTTTTACGGCGTG (SEQ ID

NO:66)

Probe CaphaP1: 5'-COCGOGGGACCCTTTAGCGA (SEQ ID

No: 67)

Conidiobolus coronatus/incongruus

Forward Primer NS92F: 5'-CACCGCCCGTCGCTAC (SEQ ID NO:68)

Reverse Primer ConiR1: 5'-TGACCAAGTTTGACCAATTTCTCTA (SEQ ID

NO:69)

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(SEQ II NO:70)

Cunninghamella elegans

Forward Primer NS92F: 5'-CACCGCCCGTCGCTAC (SEQ ID NO:71)

Reverse Primer CunR1: 5'-AATCTAGTTTGCCATAGTTCTCCTCA (SEQ

ID No:72)

Probe CunP1: 5'-TGAATGGTCATAGTGAGCATGTGGGATCTTT

(SEQ ID NO:73)

Emericella nidulans/rugulosa/quadrilineata

Forward Primer AversF1: 5'-CAACCTCCCACCCGTGAC (SEQ ID

NC:74)

Reverse Primer AniduR1: 5'-CATTGTTGAAAGTTTTGACTGATTTGT

(SEQ ID NO:75)

Probe versF1: 5'-AGACTGCATCACTCTCAGGCATGAAGTTCAG

(SEQ ID NO:76)

Eurotium amstelodami/chevalieri/herbariorum/rubrum/repens

Forward Primer EamstF1: 5'-GTGGCGGCACCATGTCT (SEQ ID

NO:77)

Reverse Primer EamstR1: 5'-CTGGTTAAAAAGATTGGTTGCGA (SEQ

ID NO:78)

Probe EamstP1: 5'-CAGCTGGACCTACGGGAGCGGG (SEQ ID

NO:79)

Epicoccum nigrum

Forward Primer EnigrF1: 5'-TTGTAGACTTCGGTCTGCTACCTCTT (SEQ ID

NO:80a

Reverse Arimer Emigral: 8'-TGCARCTGCARASSUTTTGART (SEQ ID

NO:81)

Probe EnigrP1: 5'-CATGTCTTTTGAGTACCTTCGTTTCCTCGGC

(SEQ ID NO:82)

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Reverse Primer GeoR1: 5'-TTGATTOGAAATTTTAGAAGAGCAAA (SRQ

ID NC:84)

Probe GeoF1: 5'-CAATTCCAAGAGAGAACAACGCTCAAACAAG

(SEQ ID NO:85)

Geotrichum candidum

Forward Primer NS91F: 5'-CACCGCCCCTCGCTAC (SEQ ID NO:86)

Reverse Primer GoandR1: 5'-AGAAAAGTTGCCCTCTCCAGTT (SEQ ID

110:67)

Probe Geoff: E'-TCAATCCGGAAGCCTCACTAAGCCAT1 (SEQ

ID NO:88)

Geotrichum klebahnii

Forward Primer NS91F: 5'-CACCGCCCGTCGCTAC (SEQ ID NO:89)

Reverse Primer GklabR1: 5'-AAAAGTGGCCCTCTGCTGC (SEQ ID

NO:90)

Probe GeoF2: 5'-TCAATCCGGAAGCCTCACTAAGCCATT (SEQ

ID NO:91)

Memnoniella echinata

Forward Primer StadF4 5'-TOCCAAAOCCTTATGTGAACC (SEC

ID NO:92)

Reverse Primer MemF1: 5'-TGTTTATACCACTCAGACGATACTCAAGT

(SEQ ID NO:93)

Probe MemF1: 5'-CTCGGGGCCCGGAGTCAGGC (SEQ ID

150: 24)

Mortierella polycephala/wolfii

Forward Primer MS91F: 5'-CACCGCCCGTCGCTAC (SEQ ID NO:95)

Revense Primer : MortEl: 61-76ACORAGTTTGGATTACTTTTGAG (SEC

11 1.0150.

Probe MortPl: 5'-CTTAGTGAGGCTTTCGGATTGGATCTAGGCA

(SEC ID NC:97)

Mucor mucedo

Forward Primer NS92F: 5'-CACCGCCCGTCGCTAC (SEQ ID NO:98)

Reverse Primer MmuceF.1: 5'-CTAAATAATCTAGTTTGCCATAGTTTTCG (SEQ ID NO:99)

Probe MucPl: 5'-CCGATTGAATGGTTATAGTGAGCATATGGGATC (SEQ ID NO:100)

Mucor

amphibiorum/circinclloides/heimalis/indicus/mucedo/racemosus/ramosissimus and Rhizopus azygosporus/homothalicus/microsporus/oligosporus/oryzae

Forward Primer NS92F: 5'-CACCGCCCGTCGCTAC (SEQ ID NO:101)

Reverse Primer MucR1-1: 5'-CCTAGTTTGCCATAGTTCTCAGCAG (SEQ ID NO:102)

Probe MucP1: 5'-CCGATTGAATGGTTATAGTGAGCATATGGGATC (SEQ ID NO:103)

Myrothecium verrucaria/roridum

Forward Primer MyroF1: 5'-AGTTTACAAACTCCCAAACCCTTT (SEQ

ID NC:104)

Reverse Primer MyroR1: 5'-GTGTCACTCAGAGGAGAAAACCA (SEQ ID

No:105)

Frohe MyroF1: 51-0G00TGGTT00GGG000 (SEQ ID

NÖ:106,

Paecilomyces lilacinus

Forward Frimer PlilaFl: 5'-CCCACTGTGAACCTTACCTCAG (SEQ ID

No:107)

Services and the services of the services and the services of the services of

Paecilomyces varietii

NO:110)

Forward Primer FvariF2: 5'-CGAAGACCCCTGGAACG (SEQ ID

NO:111)

Reverse Primer Frankl: 5'-GTTGTTGAAAGTTTTAATTGATTGT

(SEQ ID NO:112)

Probe FyariP1: 5'-CTCAGACGGCAACCTTCCAGGCA (SEQ

ID NO:113)

Penicillium

aurantiogriseum/polonicum/viridicatum/freii/verrucosum*/
hirsutum

Forward Primer PauraF1: 5'-CGGGCCCGCCTTTAC (SEQ ID NO:114)

Reverse Primer PauraR1-1: 5'-

GAAAGTITTAAATAATTTATATTTTCACTCAGAGTT (SEQ

ID NO:115)

Probe PenP2: 5'-CGCGCCGCCGAAGACA (SEQ ID NO:116)

Penicillium aurantiogrıseum/polonicum/viridicatum/freii

Forward Primer PauraF2: 5'-ACCGAGTGAGGGCCCTT (SEQ ID

NO:117)

Reverse Primer FauraR6: 5'-CCGGGGGGCCAGTA (SEQ ID NO:118)

Probe FenP3: 5'-TCCAACCTCCCACCGTGTTTATTT (SEQ

ID NO:119)

Penicillium brevicompactum*/alberechii

Forward Frimer PhrevF1: 5'-CCTTGTTGCTTCGGCGA (SEQ ID

NO:120)

Reverse Primer PrievR2: 5'-TCAGACTACAATCTTCAGACAGAGTTCTAA

(SEQ ID NO:121)

FireVF1: 5'-CCIGCCTTTTGGCTGCCGGG (SEQ ID

NO:122)

Penicillium

chrysogenum/griseofulvum/glandicola/coprophilum/expansum and Eupenicillium crustaceum/egyptiacum

Forward Primer

PthryF1: 5'-CGGGCCCGCCTTAAC (SEQ ID

NO:123)

Reverse Primer

PchryR1-1: 5'-

GAAAGTTTTAAATAATTTATATTTTCACTCAGAGTA (SEQ

ID

NO:124)

Reverse Primer

PchryR2-1: 5'-

GPAAGTTTTAAATAATTTATATTTTCACTCAGACCA (SEQ

ID NO:125)

Probe

PenPa: 5'-CGCGCCGCCGAAGACA (SEQ ID

NO:126)

Penicillium citrinum/sartoryi/westlingi

Forward Primer PcitrF1: 5'-CCGTGTTGCCCGAACCTA (SEQ ID NO:127)

Reverse Primer PcitrR1: 5'-TTGTTGAAAGTTTTAACTAATTTCGTTATAG

(SEQ ID NO:128)

Probe

PcitrP2: 5'-CCCCTGAACGCTGTCTGAAGTTGCA (SEQ ID

NO:129)

Penicillium corylophilum

Forward Primer ProryF1: 5'-STCCAACCTCCCACCCA (SEQ ID

NO:130)

Reverse Primer FooryR3-1: 5'-GCTCAGACTGCAATCTTCAGACTGT

(SEQ ID NO:131)

From PooryF1: 5'-01G000T0TG000C0G0S (SEQ ID

ND:132/

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Reverse Primer PdecuR3: 5'-AAAAGATTGATGTTCGGCAG (SEQ ID

NC:134)

Probe PdecuF2: 5'-CGCCGGCCGGACCTACAGAG (SEQ ID

NC:135)

Penicillium echinulatum/sclitum/camembertii/commune/crustosum

Forward Primer PohryF1: 5'-OGGGGGGGGGGGGGGTTAAG (SEQ ID MO:136)

Reverse Primer PauraR1-1: 5'-

GAAAGITTTAAATAATTTATATTTTCACTCAGAGTT (SEQ

ID MC:1371

Probe PenF2: 5'-CGCGCCGGCGGAAGACA (SEQ ID NO:138)

Penicillium expansum/coprophilum

Forward Primer PauraF2: 5'-ACCGAGTGAGGGCCCTT (SEQ ID

NO:139)

Reverse Primer PchryE6: 5'-CCCGGCGGCGCCAGTT (SEQ ID NO:140)

Probe PenP3: 5'-TOCAACCTCCCACCCGTGTTTATTT (SEQ

ID NO:141)

Penicillium fellutanum/charlesii

Forward Primer PfellF1: 5'-AACCTCCCACCGTGTATACTTA (SEQ

ID NO:142)

Reverse Primer PfellF1: 5'-CTTATCGCTCAGACTGCAAGGTA (SEQ

ID NO:143)

Probe PfellF1: OGGTTGCCCCCCGGGG (SEQ ID NO:144)

Fenicillium janthinellum/raperi

Forward Primer PjantF2: 5'-000ACCOGTGTTTATCATACCTA (SEQ

ID NO:145)

Reverse Primer PjantE2: 5'-TTSAAASTTTTAACTGATTTAGCTAATCG

'SEQ ID NO:148'

Probe PjantP2: 5'-TGCAATCITCAGACAGCGTTCAGGG (SEO

ID NO:147)

Penicillium madriti/gladioli

Forward Primer PauraF1: 5'-CGGGCCCGCCTTTAC (SEQ ID NO:148)

Reverse Primer PchryR1-1: 5'-

GAAAGTTTTAAATAATTTATATTTTCACTCAGAGTA (SEQ

ID No:149)

Reverse Primer PchryR2-1: 5'-

GAAAGTTTTAAATAATTTATATTTTCACTCAGACCA (SEQ

ID No:150)

Probe PenP2: 5'-CGCGCCGCCGAAGACA (SEQ ID NO:151)

Penicillium oxalicum

Forward Primer PoxalF1: 5'-GGGCCCGCCTCACG (SEQ ID NO:152)

Reverse Primer Poxalk1: 5'-GTTGTTGAAAGTTTTAACTGATTTAGTCAAGTA

(SEQ ID NO:153)

Probe FoxalP1: 5'-ACAAGAGTTCGTTTGTGTGTCTTCGGCG (SEQ

ID No:154)

Penicillium roquefortii

Forward Primer PchryF1: 5'-OGGGCCCGCCTTAAC (SEQ ID NO:155)

Reverse Primer FroguE2: 5'-TTAAATAATTTATATTTGTTCTCAGACTGCAT

(SEQ ID NO:156)

Probe PenP2: 5'-CGCGCCCGCCGAAGACA (SEQ ID NO:157)

Penicillium simplicissimum/cchrochloron

Forward Frimer TsimpPU-1: 5'-AACCTCCCACCCGTGTTGATT (SEQ ID NO:158;

Reverse Primer PsimpR2-1: 5'-GAGATCCGTTGTTGAAAGTTTTATCTG (SEQ ID No:159)

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Forward Primer PspinF1: 5'-GTACCTTGTTGCTTCGGTGC (SEQ ID NC:162)

Reverse Frimer PspinR1: 5'-CGTTGTTGAAAGTTTTAACTTATTTAGTTTAT (SEQ ID NO:163)

Frobe FspinF1: 5'-TCCGCGCGCACCGGAG (SEQ ID NO:164)

Rhizomucor miehei/pusillus/variabilıs

Forward Primer NE92F: 5'-CACCGCCCGTCGCTAC (SEQ ID NO:165)

Reverse Primer FmucR1: 5'-GTAGTTTGCCATAGTTCGGCTA (SEQ ID NO:166)

Frobe FmucP1: 5'-TTGAATGGCTATAGTGAGCATATGGGAGGCT (SEQ ID NO:167)

Rhizopus stolonifer

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Forward Primer NS92F: 5'-CACCGCCGTCGCTAC (SEQ ID NO:168)

Reverse Primer RstolF1: 5'-GCTTAGTTTGCCATAGTTCTCTAACAA (SEQ ID NO:169)

Frome MucPl: 5'-CCGATTGAATGGTTATAGTGAGCATATGGGATC SEQ ID NC:170)

Scopulariopsis asperula

Forward Primer SCbrvF1: 5'-CCCCTGCGTAGTAGATCCTACAT (SEQ

ID NO:171)

Reverse Primer SCasprR1: 5'-TCJGAGGTCAAACCATGAGTAA (SEQ

ID NO:172)

Probe Scopp1: 5'-TOGCATOGGGTCCOGGCG (SEQ ID

NO:173)

Scopulariopsis brevicaulis/fusca

Forward Primer SCbrvFl: 5'-CCCCTGCGTASTAGATGCTACAT (SEC

Reverse Primer SCbrvR1: 5'-TCCGAGGTCAAACCATGAAATA (SEQ ID

NO:175)

Froks SchpF1: P:+T030AC0393T0009979 SE; II

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Scopulariopsis brumptii

Forward Primer SCbrmF1: E'-CCCCTGCGTAGTAGTAAAACCA (SEQ ID

NO:177)

Reverse Primer SCbrmR1: 5'-CCGAGSTCAAACATCTTTGG (SEQ ID

No:178)

Probe ScopP1: 5'-TCGCATCGGGTCCCGGCG (SEQ ID

No:179)

Scopulariopsis chartarum

Forward Primer SIChrFl: 5'-0000CTGCGTAGTAGTAAAGC (SEO ID

NO:180)

Reverse Primer SochrR1: 5'-Toogaggtcaaaccatcaag (SEQ ID

NO:181)

Probe ScopP1: 5'-TCGCATCGGGTCCCGGCG (SEQ ID

No:182)

Scopulariopsis sphaerospora

Forward Primer SCsphF1: 5'-CCCCCTGCGTAGTAGTTTACAA (SEQ ID

NO:183)

Reverse Primer SCsphR1: 5'-CCGAGGTCAAACCATCAAAAG (SEQ ID

NO:184)

Frobe ScopP1: 5'-TCGCATCGGGTCCCGGCG (SEQ ID

No:185)

Stachybotrys chartarum

Forward Frimer StacF4 TOTOMARGOOTTATGUGAAGG 18EQ

ID NC:186)

. .

Reverse Primer StacR5 GTTTGCCACTCAGAGAATACTGAAA

(SEQ ID NO:187)

Frobe StadP2 CTGCGCCCGGATCCASGC (SFO ID

Reverse Primer TasprR2-1: 5'-GGACTACAGAAAGAGTTTGGTTGCTT

(SEQ ID NO:190)

Probe TridP1: 5'-CCAAACTGTTGCCTCGGCGGG (SEQ ID

NO:191)

Trichoderma asperellum/hamatum/viride*

Forward Primer TasprF1: 5'-CCCAAACCCAATGTGAACGT (SEQ ID

ND:192)

Reverse Primer TasprF1: 5'-TTTGCTCAGAGCTGTAAGAAATACG (SEQ

ID NO:193)

Probe TridP1: 5'-CCAAACTGTTGCCTTGGGGGG (SEQ ID

ND:194)

Trichoderma harzianum

Forward Primer TharzF1: 5'-TTGCCTCGGCGGGAT (SEQ ID

NO:195)

Reverse Primer ThanzF1: 5'-ATTTTCGAAACGCCTACGAGA (SEQ ID

110:196)

Probe TharzP1: 5'-CTGCCCGGGGTGCGTCG (SEQ ID

NO:197)

Trichoderma longibrachiatum/citroviride

Forward Primer TlongF1: 5'-TGCCTCGSCGGGATTC (SEQ ID

110:198)

Reverse Primer TlongR1: 5'-CGAGAAAGGCTCAGAGCAAAAAT (SEQ

ID NO:199)

Probe TlongPl: 5'-TOGCAGOCOCGGATCCCA (SEQ ID

110:200)

Trichoderma viride*/atroviride/koningi:

Forward Primer TviriF1: 5'-CCCAAACCCAATGTGAACCA (SEQ ID

NC:201)

Reverse Frimer TwisER1: 5'-TCCGGGGRGGGGACTRONG (SEQ ID

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Frobe TridP1: 5'-CCAAACTGTTGCCTCGGCGGG (SEQ ID

NO:203)

Ulocladium atrum/chartarum

Forward Primer UatrmF1: 5'-GCGGGCTGGCATCCTT (SEQ ID NO:204)

Reverse Frimer UatrmR1: 5'-TTGTCCTATGGTGGCGAA (SEQ ID NO:205)

Probe UloP1: 5'-TGAATTATTCACCCGTGTCTTTTGCGTACTTCT

(SEQ ID NO:206)

Ulocladium botrytis

Forward Primer UpotrF1: 5'-CCCCCAGCAGTGCGTT (SEQ ID NO:207)

Reverse Primer UbotrF1: 5'-CTGATTGCAATTACAAAAGGTTTATG (SEQ ID

(302:CM

Probe UloP1: 5'-TGAATTATTCACCCGTGTCTTTTGCGTACTTCT

(SEQ ID NO:209)

Wallemia sebi

WsebiF1: 5'-GGCTTAGTGAATCCTTCGGAG (SEQ ID NO:210)

WsebiR1: 5'-GTTTACCCAACTTTGCAGTCCA (SEQ ID NO:211)

WsebiP1: 5'-TGTGCCGTTGCCGGCTCAAATAG (SEQ ID NO:212)

Universal Fungal

ASSAY 1

Forward Primer 5.8F1: 5'-AACTITCAACAACGGATCTCTTGG (SEQ ID

NO:213)

Roverse Primer 5.8R1: 5'-GCGTTCAAAGACTCGATGATTCAC (SEO ID

NO:214:

23

Reverse Primer ZygR1: 5'-TAATGATCCTTCCGCAGGTTC (SEQ ID

No:217)

Probe ZvgP1: 5'-

CCTACGGAAACCTTGTTACGACTTTTACTTCCTCTAAA

(SEQ ID NO:218)

* Assay does not detect all strains of the indicated species

FUNGI

Aspergillus auricomus

Forward Primer AdirdF1: 5'-ATTACTGAGTGAGGGTCCCTCG

Reverse Primer AauriR1: 5'-GGCGGCCGCGTAAAC

Probe AdirdP1: 5'-000G00GAAGCAACAAGGTACG

Aspergillus caespitosus

Forward Primer AcaesF1: 5'-CTCCCACCCGTGAATACCTT

Reverse Primer AcaesR1: 5'-GGCTCAGACGCAACTCTACAAT

Frcbe AcaesP1: 5'-CACTGTTGCTTCGGCGAGGAGCC

Aspergillus candidus

Forward Primer AdandF1: 5'-TTACCGAGTGAGGGTTTCTCTGA

Reverse Primer AcandR1: 5'-ACAGTGTTGGGGTCTT

Probe PsimpP1: 5'-00300TCA0GGCGCGC

Aspergillus cervinus

Forward Primer AdervF1: 5'-CCACCCGTGCTATTGTACCTTT

Peverse Primer AdervR1-1: 5'-CARCPOAGROTGGLATTGPGRACtGT

Probe AfumiP2: 5'-TTCTCGGCGGGGCGCGG

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Forward Primer AclayF1: 5'-CCCGCCGTCTTCGGA

Reverse Primer AclavR1: 5'-CCGTTCTTGAAAGTTTTAACTGATTATG

Frobe AfumiP1: 5'-CCCGCCGAAGACCCCAACATG

Aspergillus flavipes

Forward Primer AflvpF1: 5'-CCACCCGTGACTACTGTACCAC

Feverse Primer AflypR1: 5'-CCGGCGGCGCCAGCTAG

Feverse Primer AflvpR2: 5'-A3GCTTTCAGAAACAGTGTTC3

Frobe AspP1: 5'-TTGCTTCGGCGGGCCC

Aspergillus niveus

Forward Primer AniveF1: 5'-ACCCGTGCCTATTGTACCCT

Fleverse Primer AniveR1: 5'-TGCAAACAATCACACTCAGACAC

Frobe AspP1: 5'-TTGCTTCGGCGGGCCC

Aspergillus ochraceus

Forward Primer AcchrF1: 5'-AACCTCCCACCGTGTATACC

Figure Primer AochrR2-1: 5'-CGGCGAGCGCTGTtCC

Probe AcchrPl: 5'-ACCTTGTTGGTTGGGCGAGCCC

Aspergillus ostianus

Forward Primer AochrF1: 5'-AACCTCCCACCGGTGTATACC

Reverse Primer AbstiR1-1: 5'-05G0GAG0G0TGTtGT

Am orgalilus parademus

Reverse Primer ApardR1-1: 5'-GACTGCAACTTCATACAGAGTTGGT

Probe PenP2: 5'-CGCGCCGGCGAAGACA

Aspergillus penicillicides

Forward Primer ApeniF2: 5'-0GCCGGAGACCTCAACC

Reverse Primer ApeniR2: 5'-TCCGTTGTTGAAAGTTTTAACGA

Probe ApeniP2: 5'-

TGAACACTGTCT BAA BETTGCAGTCTGAGTATG

Aspergillus sclerotiorum

Forward Primer AcircF1: 5'-ATTACTGAGTGAGGGTCCCTCG

Reverse Primer AsclrR1: 5'-CCTAGGGAGGGGGTTTGA

Probe AcircF1: 5'-CCCGCCGAAGCAACAAGGTACG

Aspergillus sydowii

Forward Primer AsydoFl-1: 5'-CAACCTCCCACCCGaGAA

Reverse Primer versR1: 5'-CCATTGTTGAAAGTTTTGACTGATTTTA

Probe versP1: 5'-AGACTGCATCAGTCTCAGGCATGAAGTTCAG

Aspergillus unguis

Forward Frimer AunguFl: 5'-CAACOTCOCAGGCTTGAATAGT

Reverse Primer AunguR1: 5'-TCACTCTCAGGCATGAAGTTCAG

Probe AcaesP1: 5'-CACTGTTGCTTCGGCGAGGAGCC

Aspergillus wentii

Forward Frimer AwentF1: 5'-CATTACCGAGTGAGGACCTAACC

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Probe AdirdP1: 5'-CCCGCCGAAGCAACAAGGTACG

Candida albicans

Forward Primer CalbF1: 5'-CTTGGTATTTTGCATGTTGCTCTC

Feverse Primer CalbR1: 5'-GTCAGAGGCTATAACACACAGCAG

Frobe CalbF1: 5'-TTTACCGGGCCAGCATCGGTTT

Candida dubliniensis

Forward Primer CdubF1: 5'-AGATCAGACTTGGTATTTTGCAAGTTA

Fleverse Primer | CdubFl: 5'-TAGGCTGGCAGTATCGTCAGA

Frobe CdubF1: 5'-TTTACCGGGCCAGCATCGGTTT

Candida (Pichia) guilliermondii

Forward Primer CguiF1: 5'-CCTTCGTGGCGGGGTG

Faverse Primer CguiF1: 5'-GCAGGCAGCATCAACGC

Frobe CguiP1: 5'-CCGCAGCTTATCGGGCCAGC

Candida haemulonii

Forward Frimor ChaeF1: 5'-GGAGUGACAAGGAGCAGTC

Reverse Primer ChaeR1: 5'-AGGAGGCAGAAAGGAAGAGG

Probe ChaeP1: 5'-ATSTASTACAGCCCTCTGGGCTGTGCA

Probe Cha2P1: 5'-AAGIGGGAGCTGATGTAGCAACCCCC

Candida krusei

Forward Primer CkruF1: 5'-CTCAGATTTGAAATCGTGCTTTG

Reverse Primer CkruF.1: 5'-GGGGCTCTCACCCTCCTG

Probe CkruF1: 5'-CACGAGTTGTAGATTGCAGGTTGGAGTCTG

Candida lipolytica

Forward Primer ClipF1: 5'-TAGCGAGAGGGGTTACAAATG

Peverse Primer ClipF.1: 5'-CGTCGGTGGCAGTGTGGA

Frobe ClipF1: 5'-CCTTCGGGGGTTCTCCCCTAACC

Candida lusitaniae

Forward Primer ClusF1: 5'-GGGCCAGCGTCAAATAAAC

Fleverse Primer ClusF1: 5'-CGCAGGCCTCAAACAACA

Frobe ClusP1: 5'-AGAATGTGGCGTGCCTTCG

Candida maltosa

Forward Primer CmalF1: 5'-3GCCAGCATCAGTFTGGAC

Fleverse Primer | CmalFil: 5'-TOTAGACTGGCAGTATCGACAGTG

Frobe CmalP1: 5'-IAGGACAATTGCGGTGGAATGTGGC

Candida parapsilosis

Forward Primer CparF1: 5'-GATCAGACTTGGTATTTTGTATGTTACTCTC

Probe CparP1: 5'-CCTCTACAGTTTACCGGGCCAGCATCA

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Candida sojae

Forward Primer CsojF1: 5'-CGGTTGTGTGTTATAGCCTTCGTA

Reverse Primer CsojR1: 5'-ATCATTATGCCAACATCCTAGGTAAT

Probe CtropP2: 5'-CGCAGTCCTCAGTCTAGGCTGGCAG

Candida tropicalis

Forward Primer CtropF1: 5'-GCGGTAGGAGAATTGCGTT

Reverse Primer CtropR2: 5'-TCATTATGCCAACATCCTAGGTTTA

Probe CtropP2: 5'-CGCAGTCCTAGGCTGGCAG

Candida viswanathii

Forward Primer CvisF1: 5'-CGGCAGGACAATCGCGT

Reverse Primer CvisR1: 5'-TCTAGGCTGGCAGTATCCACG

Probe CvisP1: 5'-AATGTGGCACGGCCTCGGC

Candida zeylanoides

Forward Primer Czey F1: 5'-GTTGTAATTTGAAGAAGGTAACTTTGATT

Reverse Primer Czey R1: 5'-GACTCTTCGAAAGCACTTTACATGG

Probe Chey P1: 51-00TTGGAACAGGGGTCACAGAGGGGT

Emericella (Aspergillus) nidulans/rugulosa/quadrilineata

Forward Primer AversF1: 5'-CAACCTCCCACCCGTGAC

Feverse Primer AniduR1-1: 5'-CCATTGTTSAAAGTTTTSACTGATaTGT

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Forward Primer GklebF1: 5'-GGGGGACTTTTCCGGC

Reverse Primer GklebR2: 5'-TGGCACAAATTCTCCTCTAATTTATTTA

Probe GklebP1: 5'-

AAGCTAGTCAAACTTGGTCATTTAGAGGAAGTAAAAGTC

Penicillium aethiopicum

Forward Primer PaethF1-1: 5'-0GGGGGGGCTCtCGCT

Reverse Primer PohryR1-1: 5'-

GAAAGTTTTAAATAATTTATATTTTCACTCAGAgTA

Probe

PenP2: 5'-CGCGCCCGCCGAAGACA

Penicillium atramentosum

Forward Primer PgrisF1-1: 5'-ACCTGCGGAAGGATCATTtCT

Reverse Primer PatraR1: 5'-CCCCGGCGGCCATA

Probe

PenP3: 5'-TCCAACCTCCCACCCSTGTTTATTT

Penicillium aurantiogriseum

Forward Primer PauraF3: 5'-CGCCGGGGGGGCTTC

Reverse Primer PauraR1-1: 5'-

GAAAGTTTTAAATAATTTATATTTTCACTCAGAgTT

Probe

PenF2: 5'-030G000G00GAAGACA

Penicillium aurantiogriseum/polonicum/viridicatum/freii

Forward Primer PexpaF1-1: 5'-TTACOGAGTGAGGGCCGTT

Reverse Framer FauraR6: 5'-000gg0gg0CAgTA

Probe

PenP3: 5'-TOGAACCTOGCACCGGTGTTTATTT

Fenicillium canescens

Forward Primer FcaneF1: 5'-TTACCGAGCGAGAATTCTCTGA

Reverse Frimer PcaneR1: 5'-AGACTGCAATTTTCATACAGAGTTCA

Frobe PsimpP1: 5'-CCGCCTCACGGCCGCC

Penicillium citreonigrum

Forward Primer PoteoF1-1: 5'-TGTTGGGCTCCGTCCTCtTC

Feverse Primer PoteoR1-1: 5'-CGGCCGGGCCTtCAG

Probe PenP7: 5'-CCGAAAGGCAGCGGCGC

Penicillium coprophilum

Forward Primer ProprF1-1: 5'-GGGTCCAACCTCCCACtCA

Reverse Primer PchryR1-1: 5'-

GAAAGTTTTAAATAATTTATATTTTCACTCAGAgTA

Probe PenP1: 5'-CGCCTTAACTGGCCGCCGG

Penicillium crustosum

Forward Primer PcrusF1: 5'-CGCCGGGGGGCTTA

Reverse Prime: FauraR1-1: 5'-

GRAAGTTTRAATAATTATATTTTCACTCAGAGTT

Probe PenF2: 5'-CGCGCCGCCGAAGACA

Penicillium digitatum

Penicillium expansum

(assay 1)

Forward Primer PexpaF1-1: 5'-TTACCGAGTGAGGGCCgTT

Reverse Primer FexpaR2-1: 5'-GCCCGCCGAAGCtACG

Probe FenP3: 5'-TCCAACCTCCCACCCGTGTTTATTT

(assay 2)

Forward Primer PexpaF2-1: 5'-TCCCACCCGTGTTTATTTACaTC

Reverse Primer FexpaR1: 5'-TCACTCAGACGACAATCTTCAGG

Probe FenF1: 5'-CGCCTTAACTGGCCGCCGG

Penicillium freeii

Reverse Primer FauraR1-1: 5'-

GAAAGTTTTAAATAATTTATATTTTCACTCAGAgTT

Prohe FenP2: 5'-CGCGCCGCCGAAGACA

Penicillium glandicola

Forward Primer FglanF1-1: 5'-CCGGGGGGGCTTtCGT

Reverse Primer FchryR1: 5'-

GAAAGTTTTAAATAATTTATATTTTCACTCAGACTA

Probe PenP2: 5'-CGCGCCGGGGAAGACA

Penicillium griseofulvum

Forward Primer FgrisF1-1: 5'-ACCTGCGGAAGGATCATTtCT

Reverse Primer FonryR6: 51-000990G900ASTT

Probe PenP3: 5'-TCCAACCTCCCACCCGTGTTTATTT

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Penicillium hirsutum*

Forward Primer PhirsF1-1: 5'-GCCGGGGGGCTCAtA

Feverse Primer PauraF.1-1: 5'-

GAAAGTTTTAAATAATTTAIATTTTCACTCAGAgTT

Frobe PenP2: 5'-CGCGCCGCCGAAGACA

Penicillium implicatum

Forward Primer PimplF1: 5'-GCCGAAGACCCCCCTGT

Reverse Primer PimplF.1: 5'CGTTGTTGAAAGTTTTGACTGATTGT

Frobe PimplP1: 5'-AACGCTGTCTGAAGCTTGCAGTCTGAGC

Penicillium islandicum

Forward Primer PislaF1: 5'-CGAGTGCGGGTTCGACA

Reverse Primer PislaR1: 5'-GGCAACGCGGTAACGGTAG

Probe PislaP1: 5'-AGCCCAACCTCCCACCCGTG

Penicillium italicum

Forward Primer PitalF1-1: 5'-CTCCCACCCGTGTTTATTTAtCA

Feverse Frimer FempaR1: E'-TORUTCAGAGGAGGACGACTCAGG

Revense Brimer | FempaRC+1: E'-ICACTCAGACGACGAUCTTCEGG

Probe FemF1: (+) 5'-0300TTAACTGGCCGCCGG

Penicillium melinii

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Penicillium miczynskii

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Forward Primer PmiczF1-1: 5'-GTGTTTAACGAACCTTGTTGCaTT

PmiczR1-1: 5'-CTCAGACTGCATACTTCAGACaGA Feverse Primer

Frobe PsimpF1: 5'-COGOCTCACGGCCGCC

Fenicillium olsonii

Forward Primer PclsnF1: 5'-GGCGAGCCTGCCTTCG

Fleverse Primer PenR2: 5'-GATCCGTTGTTGAAAGTTTTAAATAATTTATA

Frobe PolsnF2: 5'-TCCGCGCTCGCCGGAGAC

Penicillium purpurogenum

Forward Primer PpurpF1: 5'-ASGATCATTACTGAGTGCGGA

Reverse Primer PpurpF.1: 5'-GCCAAAGCAACAGGGTATTC

Frobe PpurpP1: 5'-CCCTCGCGGGTCCAACCTCC

Penicillium raistrickii

Forward Primer PgrisF1-1: 5'-ACCTGCGGAAGGATCATTtCT

Fleverse Primer PraisF.1: 5'-CCCGGCGGCGGCCAGAC

PenP3: 5'-TCCAACCTCCCACCCGTGTTTATTT Probe

Penicillium restrictum

Forward Frimer PrestF1-1: 5'-CACGGCTTGTGTGTTGGGtCT

Reverse Primer PrestR1-1: 5'-CGGCCGGGCCTaCAA

Probe PenP7: 5'-CCGAARGGCAGCGGCGGC

Penicillium sclerotiorum

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Forward Primer PsclrF1: 5'-TTCCCCCGGGAACAGG

Peverse Frimer FsclrR1: 5'-GCCCCATACGCTCGAGGAT

Frobe PsclrPl: 5'-CCGAAAGGCAGTGGCGGCAC

Penicillium simplicissimum/ochrochloron

Forward Frimer PsimpF2-1: 5'-CGCCGAAGACACCATTGAtCT

Feverse Primer PsimpR4-1: 5'-CTGAATTCTGCAATTCACATaACG

Probe PsimpP2: 5'-TGTCTGAAGATTGCAGTCTGAGCGATTAGC

Penicillium variabile

Forward Frimer PvarbF1: 5'-GCCGGGGGGCTTCT

Fleverse Frimer PvarbF1: 5'-TCTCACTCAGACTCACTGTTCAGG

Probe PvarbP1: 5'-AGGGTTCTAGGGTGCTTCGGCGG

Penicillium verrucosum*

Forward Frimer PverrF2: 5'-CGGGCCCGCCTTTG

PauraR1: 5'-

GAAAGTTTTAAATAATTTATATTTTCACTCAGACTT

Probe PenPO: 5'-0603000300GAAGACA

Penicillium waksmanii

Forward Primer F waksF1-1: 5'-GTGTTTAACGAACCTTGTTGCATC

Reverse Primer P waksF1-1: 5'-CTICAGAGAGGGTTGAGAGGTAG

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Forward Primer UatrmF2: 5'-CGGGCTGGCATCCTTC

Reverse Primer UatrmR2: 5'-GTGAITGCAATTACAAAAGGTTTATG

Probe UloP1: 5'-

TGAATTATTCACCOGTGTCTTTTGCGTACTTCT

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Ulocladium chartarum

Forward Primer UpharFl-1: 5'-AGCGGGCTGGAATCCaTT

Fleverse Primer UcharR1-1: 5'-CTGATTGCAATTACAAAAGGTTgAAT

Frobe UloP1: 5'-

TGAATTATTCACCCGTGTCTTTTTGCGTACTTCT

Universal Fungal

Forward Primer 5.8F1-1: 5'-AACTTTCAACAACGGATCTCTTG

F.everse Primer 5.8R1-1: 5'-CGTTCAAAGACTCGATGATTCAC

Probe 5.8P1: 5'-CATCGATGAAGAACGCAGCGAAATGC

BACTERIA

Legionella maceachernii

Reverse Primer PmaceR1: 5'-JACJACCCTCTCCTATACTCTTAGTCCAG

Frobe LmicdP1:

5'-AGTCTTATCTGACCACCCTAGGTTGAGCCCA

Legionella micdadei

Forward Primer LmicdF1: 5'-GGTGGTTTTATAAGTTATCTGTGAAATTC

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Reverse Primer PmicdR1: 5'-CACTACCCTCTCCTATACTCAAAGTCTC

Frobe LmicdF1:

5'-AGTCTTATCTGACCACCCTAGGTTGAGCCCA

Legionella pneumophila

(Type1)

Forward Primer LpneuF1: 5'-CGGAATTACTGGGCGTAAAGG

Reverse Primer PpneuF.1: 5'-GAGTCAACCAGTATTATCTGACCGT

Frobe LpneuF1:

5'-AAGCCCAGGAATTTCACAGATAACTTAATCAACCA

(Type 2)

Forward Primer LpneuF3: 5'-CCCAGCTTTCGTCCTCAGAC

Reverse Primer LpneuR2: 5'-AGTCGAACGGCAGCATTG

Frcbe LpneuP2: 5'-TGCTAGACAGATGGCGAGTGGCGA

Legionella sainthelensi/cincinnatiensis

Forward Primer LsainF1: 5'-CGTAGGAATATGCCTTGAAGACT

Reverse Primer PsainF1: 5'-AAGGTCCCCAGCTTTCGT

Prote LsainPl:

E'-AGACATURTOCGGTATTARUTTGAGTTTCCC

Aercmonas hydrophila

Forward Primer AbydF1: 5'-TGCCGCGTGTGTGAAGAA

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Aeromonas media/eucrenophila

Forward Primer AmedF1: 5'-ATGCCGCGTGTGTGAAGA

Reverse Primer AmedR1: 5'-CGAGTAACGTCACAGCTGATG

Frobe AmedP1: 5'-AAGCACTTTCAGCGAGGAAAGGTTG

Aeromonas schubertii

Forward Primer AschF1: 5'-AGCGAGGAGGAAAGGTTGGT

Fleverse Primer AschEl: 5'-GGAGTTAGCCGGTGCTTCTTC

Probe AschP1: 5'-TGCGAGTAACGTCACAGCTGGCAGGTAT

Aeromonas veroni

Forward Primer AverF2: 5'-AGCGAGGAGGAAAGGTTGGTAG

F.everse Primer AverR2: 5'-CGGAGTTAGCCGGTGCTTC

Probe AverF2: 5'-TAATAACTGCCAGCTGTGGACGTTACTCGCA

Aeromonas caviae, trota, jandaei

Forward Primer AverF2: 5'-AGCGAGGAGGAAAGGTTGGTAG

Reverse Primer AcavE1: 5'-03GAGTTAGCCGGTGCTTC

Probe AcavP1: 5'-TCTGCGAGTAACGTCACAGCCAGCAGATA

Aeromonas all species

Forward Primer AuniF1: 5'-CAGGGCTACACACGTGCTACA

Reverse Primer AuniR1: 51-GGGATTGGGTCRGTRTGGGT

Probe AuniP1: 5'-TGGCGCGTACAGAGGGCTGCA

Table 2. List of Bacterial Primers and Probes

Escherichia coli					
Forward Primer	uidAF1: No:219)	5'-GGGCAGGCCAGCGTATC (SEQ ID			
Reverse Primer	uidAR1: NO:220)	5'-CCCACACTTTGCCGTAATGA (SEQ ID			
Reverse Primer	uidAF.2: No:221)	5'-CGTACACTTTTCCCGGCAAT (SEQ ID			
Probe	uidAF1: NO:222)	5'-TGCTGCGTTTCGATGCGGTCA (SEQ ID			
Helicobacter pylorii					
Forward Primer	HpylF1: No:223)	5'-GGGTATTGAAGCGATGTTTCCT (SEQ ID			
Reverse Primer	HpylF.1: NO:224)	5'-GCTTTTTTGCCTTCGTTGATAGT (SEQ ID			
Probe		5'-AAACTCGTAACCGTGCATACCCCTATTGAG NO:225)			

One skilled in the art will appreciate that primers and/or probes can be used which are not identical to the ones described above, as long as there is substantial similarity between the sequences. Of purposes of the present invention, "substantial similarity" means that more than 90-110% of the sequence is the same as the sequences enumerated above.

Performance of Assav

Standard procedures for the operation of the model

changed from 50 to 25 μ l. Thermal cycling conditions consisting of two min at 50° C, 10 min at 95° C, followed by 40 cycles of 15 sec at 95° C and 1 min at 60° C. Cycle threshold (C_T) determinations, i.e. non-integer calculations of the number of cycles required for reporter dye fluorescence resulting from the synthesis of PCR products to become significantly higher than background fluorescence levels were automatically performed by the instrument for each reaction using default parameters. Assays for fungal target sequences and G. candidum (reference) sequences in the same DNA samples are performed in separate reaction tubes.

Quantification of fungal target

Quantification is performed by first subtracting mean reference sequence C_T values from mean target sequence C_T values for both test samples and a pre-specified calibrator sample to obtain ΔC_T values. Calibrator sample ΔC_T values are then subtracted from ΔC_T values of the test samples to obtain $\Delta \Delta C_T$ values. Assuming an amplification efficiency of one (i.e. a doubling of the target sequence for each cycle), the ratio of target sequences in the test and calibrator samples is given by $2^{-\Delta\Delta C_T}$. (If the efficiency is less than one, then the new amplification efficiency value is used instead of 2.) For example, a ratio of 0.1, calculated in this manner, would indicate that the target sequence level in the test sample is chartenth the level found in the tallprator sample. A circulated comparison (ΔC_T) approach should allow the discrimination of 1-

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fold differences in the quantities of target sequences in different samples with 95% confidence.

Specific Examples

Example 1:

Quantitative Measurement of Stachybotrys chartarum conidia using real time detection of PCR products with the $TaqMan^{TM}$ Fluorogenic probe system

Conidial stocks of the target fungus, e.g.

Stachybotrys chartarum, and the reference target, e.g.

Geotrichum candidum, were prepared to act as calibrator and internal standard, respectively.

Genomic DNAs were extracted from 20µl conidial suspensions using a glass bead milling and glass milk adsorption method. Briefly, this method involved mixing test and reference conidia suspensions (10µl ea.) with 0.3g of acid-washed glass beads (G-1277; Sigma, St. Louis, MO, USA) and 10µl, 100µl and 300µl, respectively, of glass milk suspensions, lysis buffer and binding buffer from an Elu-Quik INA purification kit (Schleicher and Schuell, Keene, NH) in sterile 2ml conical bottom, screw cap tubes (506-636; FGC Scientifics, Gaithersburg, MD). The tubes were shaken in a mini beadbeater (Biospec Products, Bartlesville, OK) for one minute at maximum rate and FNAs were recovered in final

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The TagMan probes and primers were obtained from the custom oligonucleotide synthesis facility at PE-Applied Biosystems (Foster City, CA). TagMan probes contained a TAMRA group conjugated to their 3'-terminal nucleotide and a FAM group linked to their S'-terminal nucleotides as the quencher and reporter fluorochromes, respectively. For Geotrichum candidum, the forward primer is NS92F: 5'-CACCGCCCGTCGCTAC (SEQ ID NC:86), the reverse primer is GeandR1: 5'-AGAAAAGTTGCCCTCTCCAGTT (SEQ ID NO:87), and the probe is GeoP2: 5'-TCAATCCGGAAGCCTCACTAAGCCATT (SEQ ID NO:88). For Stackybotrys chartarum, the forward primer is StacF4 5'-TCCCAACCCTTATGTGAACC (SEQ ID NO:186), the reverse primer is StacR5 5'-GTTTGCCACTCAGAGAAATACTGAAA (SEQ ID NO:187), and the probe is StacP2 5'-CTGCGCCCGGATCCAGGC (SEQ ID NO:188).

PCR reactions were prepared in 0.5ml thin-walled, optical grade PCR tubes (PE Applied Biosystems, Foster City, CA) by addition of the following components: 12.5µl of TaqMan Universal Master Mix (a 2x-concentrated, proprietary mixture of AmpliTaq Gold™ DNA polymerase, AmpErase® UNG, dNTPs with UTP, passive reference dye and optimized buffer components, PE Applied Biosystems, Foster City, CA); 2.5µl of mixture of forward and reverse primers (10nM each); 2.5µl of 400nM TaqMan probe; 2.5µl of 2mg/ml bovine serum albumin, fraction V (Sigma Chemical, St. Louis, MO) and 5µl of DNA template. Standard procedures for the operation of the model 7700, as described in the instrument's manual, were followed. This included the

use of all default program settings with the exception of reaction volume which was changed from 50 to $25\mu l$. Thermal cycling conditions consisting of two min at $50^{\circ}C$, 10 min at $95^{\circ}C$, followed by 40 cycles of 15 s at $95^{\circ}C$ and 1 min at $60^{\circ}C$. Cycle threshold (C_{T}) determinations, i.e. non-integer calculations of the number of cycles required for reporter dye fluorescence resulting from the synthesis of PCR products to become significantly higher than background fluorescence levels, were automatically performed by the instrument for each reaction using default parameters. Assays for S. Chartarum (target) sequences and G. candidum (reference) sequences in the same DNA samples were performed in separate reaction tubes.

Quantification of S. chartarum conidia using the comparative C_T method was performed by first subtracting mean reference sequence C_T values from mean target sequence C_T values for both test samples and a pre-specified calibrator sample to obtain ΔC_T values. Calibrator sample ΔC_T values are then subtracted from ΔC_T values of the test samples to obtain ΔC_T values.

Calibrator samples were DNA extracts from mixtures of approximately 2 x 10^4 S. chartarum (strain UMAH 6417) and 2 x 10^4 S. candidum schidia. Tost samples were mixed with the

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quantities of S. chartarum conidia in the calibrator samples to obtain estimates of the absolute quantities of these conidia in the test samples.

Each series of DNA extracts was also analyzed using only S. chartarum target sequence assay results. In these calculations, calibrator sample C_T values were subtracted directly from corresponding test sample C_T values to obtain $\Delta C_{T,STAE}$ values. These values were used in place of $\Delta\Delta C_T$ values to determine the ratio of target sequences in the test and calibrator samples and to quantify S. chartarum conidia in the test samples as described above.

Air sampling was performed in rooms that had previously been occupied by infants diagnosed with PH from three in the Cleveland, Ohio area. Airborne particles were recovered in sterile BioSampler® vials (SMC Inc., Eighty Four, PA) connected to an AirCon-2 High Flow Sampler pump (Gilian Instrument Co., Clearwater, FL). Air samples were taken over an eight hour time period under passive conditions (i.e. with no activity occurring in the rooms) at a flow rate of 10 liters per min, for a total collection volume of 4.8m³. Two additional air samples were taken in the same manner over a twelve hour period from the basement of a home in the Cincinnati, Ohio area that was also determined to contain extensive S. chartarum growth. One of these samples was collected under passive conditions, as described above, while the other was collected under aggressive sampling conditions

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(i.e. during and after a cleaning effort in the contaminated area).

Each of the BioSampler vessels was rinsed three times with 5ml of sterile distilled water. The pooled rinses from each vial were transferred to sterile 50ml capped test tubes (25330-50); Corning Inc., Corning, NY) and centrifuged for 15min at 1000 x g in a Sorval RC2 centrifuge using an SS-34 rotor (DuPont Instruments, Newton, CT). After carefully drawing off the top 13-14ml of the supernatants, the pelleted materials in each tube were resuspended in the remaining liquid and transferred to 2ml microfuge tubes (16-8100-27; PGC Scientific, Frederick, MD). These suspensions were centrifuged at 14,000rpm for 3min in an Eppendorf microcentrifuge (5415C; Brinkman Instruments, Westbury, NY) and the majority of the supernatants were again removed by pipetting. The pellets and small amounts of liquid remaining in each tube were adjusted to either 100 or 200µl with sterile distilled water.

Direct counts of putative S.chartarum conidia in 1901 aliquots of the recovered samples were made in a haemocytometer chamber. Separate counts of up to six aliquots of each samples were taken over the entire grid portion of the chamber and the mean counts were converted to cell

chartarum conidia were based on recognition of the characteristic size, shape and pigmentation of these conidia. Three additional 10pl aliquots of each recovered sample were mixed with *G. candidum* reference conidia and subjected to total genomic DNA extraction for subsequent analysis in the model 7700 as specified above.

Yields of target sequences extracted from these conidia samples and from calibrator samples were determined from their respective $C_{\overline{\tau}}$ results in the model 7700 and compared using both the $\Delta\Delta C_T$ (including Gestrichum reference sequence data) and $\Delta C_{T,STAC}$ (not including reference sequence data) versions of the comparative C, method. Quantities of conidia estimated from these analyses were then comparted with those determined from direct microscopic counts of the samples taken in a haemocytcmeter. As illustrated in Fig. 3, results obtained by the $\Delta C_{T,\,\text{eTAC}}$ analysis method and from direct counting showed good agreement for most the samples. In 13 of these 14 instances, the estimate of the $\Delta C_{T,STAC}$ method was within a onefold range of the direct counting result. The results further indicated that this level of presumed accuracy and precision (i.e. within a 50-200% range of direct counts) may be expected to occur in 95% of all analyses performed by the $\Delta C_{T,\,STAC}$ method. Based on comparisons of results obtained by the $\Delta\Delta C_T$ analysis method for the same samples (data not shown), it was estimated that this method would provide the same revel of accuracy in only about 70% of all analyses. Conidia from each of the

different strains examined appeared to be quantified with similar degrees of precision and accuracy using the $\Delta C_{T,STAC}$ analysis method.

The sensitivity of the TaqMan assay and the functional dynamic range of the $\Delta C_{T,STAC}$ quantification method were further examined using ten-fold serial dilutions of S. chartarum strain UMAH 6417 conidia stock suspensions as test samples. These samples contained expected quantities of cells that ranged from 2 to 2,000 based on direct counting analyses of the starting stock suspensions. As shown in Fig. 4, the results of these analyses were again in good agreement with the expected results. Five of the eight measurements gave estimates that coincided with the expected quantities of conidia in the samples within the relative errors of the analyses. The mean results of these analyzers were within a one-fold range of the expected values in all instances. In one of these two experiments, a low level of signal (equivalent to an estimated mean quantity of 0.27 conidia) was observed in the negative control samples. Farallel samples taken from one of the two dilution series of conidia (cf. Experiment 2 in Fig. 47 were also subjected to DNA extractions in the absence of Geotrichum cells. Although these extract yielded slightly lower quantitative results than those

data not shown).

method was made by analyzing particulate samples collected from the inside air of four homes with known colonization of S. chartarum. TaqMan-based results were again compared with those obtained by direct microscopic observations of the samples in a haemocytometer. As shown in Table 3, the two methods again gave similar mean determinations of the quantities of S. chartarum conidia in these samples with four of the five results agreeing within the relative errors of the TaqMan analyses. No S. chartarum conidia were found in the fifth samples by direct microscopic observation, however, this sample also appeared to approach the detection limits of the TaqMan assay with only two of the three replicate DNA extracts producing signals above background.

Example 1, Table 1. Quantification of Stachybotrys chartarum conidia recovered from indoor air samples by direct microscopic counting and the $\Delta C_{T,STAC}$ method as determined from TaqMan analysis.

Sample	Sampling	Direct count estimate	<u> AC_{T, STAC} TacMan estimate</u>		
contce	conditions*	Conidia, m ⁻¹ air	Conidia, m ^{ri} air	Relative error	
Home 1b	Passive	46°	23 ^b	7.5-69	
Home 2 ^b	Passive	15°	14°	5.2-37	
Home 3b	Passive	31°	2 6°	9.4-68	
Home 4t	Passive	O.c.	2.2°	0.3-19	
Home 4 ^b	Aggressive	5 6 0 0°	4300	2660-730	

Defined in Materials and Methods.

Example 2:

Quantification of fungus from dust using real time, fluorescent probe-based detection of PCR products

Dust samples from the home of an infant with pulmonary hemosiderosis in Cleveland, OH (Home 1) were collected using 37-mm filter cassettes, pore size 0.8 μm , as the collection device. Samples were obtained from two rooms in the basement, the living room, and the dining room. Additional dust samples were obtained in a similar manner from the basement of a home in Cincinnati, OH (Home 2) containing a significant, but localized, growth of S. chartarum as determined by surface sample analysis. One sample was taken from the floor directly beneath the area of growth, a second from another location in the same room and a third from an the specific control is the second of the se

b Located in Cleveland, Ohio.

 $^{^{\}circ}$ Value based on a total air sample volume of 4.8 m $^{\circ}$.

Located in Cincinnati, Ohio.
 Value based on a total air sample volume of 7.2m³.

Total DNAs were extracted from dust samples using glass bead milling and glass milk adsorption method. Weighed dust samples were added directly to sterile 2 ml conical bottom, screw cap tubes (506-636; PGC Scientifics, Gaithersburg, MD), containing 0.3 g of glass beads (G-1277; Sigma, St. Louis, M0) and 100 and 300 μl of lysis and binding buffer, respectively from an Elu-Quik DNA Purification Kit (Schleicher and Schuell, Keene, NH). Ten ul aliquots cf a 2 X 107 conidia/ml suspension of G. candidum in 0.5% Tween 20 were also routinely added to the tubes as a potential source of reference DNA sequences. Ten µl aliquots of S. chartarum conidia suspensions in water were also added as needed. The tubes were shaken in a mini beadbeater, (Biospec Products, Bartlesville, OH) for one minute at a maximum speed. To bind the DNA, 25 µl of Elu-Quik glass milk suspension (Schleicher and Schuell, Keene, HN) was added to the samples and the tubes were placed on a minimutating mixer (Glas-Col, Terre Haute, IN) for 20 minutes. The samples were transferred to $SPIN^{m}$ filter and catch tube assemblies (BIO 101, Vista CA) and centrifugation at 7500 X g for 1.5 min to remove binding and lysis buffers. The retained particulates, including glass milk with adsorbed nucleic acids, were washed twice in the filter cartridges with 0.5 ml Elu-Quik wash buffer and once with 0.5 ml Elu-Quik salt reduction buffer and centrifuged as above alitus egos Maan, september aproped from the glass milk particles by two successive washes with 100 µl distilled

water and collected by centrifuging the washes into clean catch tubes. Calibrator samples, used in the analytical method as standards for the quantification of S. chartarum conidia in the test samples, contained 2 \times 10⁴ S. chartarum and 2 \times 10⁵ G. candidum conidia with nc dust and DNA extractions from these samples was performed in the same manner.

PCR reactions were prepared in 0.5 ml thin-walled, optical grade FCR tubes (PE Biosystems, Foster City CA). Each reaction contained 12.5 µl of "Universal Master Mix"- a 2X concentrated, proprietary mixture of AmpliTaq Gold™ DNA polymerase, AmpErase® UNG, dNTPs, passive reference dye and optimized buffer components (PE Biosystems, Foster City CA), 0.5 µl of a mixture of forward and reverse primers at 50 mM each, 2.5 µl of 400 nM TaqMan probe (PE Biosystems, Foster City, CA), 2.5 µl of 2 mg/ml fraction V bovine serum albumin (Sigma Chemical, St. Louis, MC) and 2 µl of autoclaved water. Five µl of purified DNA extract was added to complete the 25 µl reaction mix.

The TagMan probes and primers were obtained from the custom eligenuclectide synthesis facility at FE-Applied Eicsystems (Foster City, CA). TagMan probes contained a TAMRA group conjugated to their 3'-terminal nucleotide and a FAM group linked to their 5'-terminal nucleotides as the guencher

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AGAAAAGTTGCCCTCTCCAGTT (SEQ ID NO:87), and the probe is GeoP2: 5'-TCAATCCBGAAGCCTCACTAAGCCATT (SEQ ID NO:88). For Stachybotrys chartarum, the forward primer is StacF4 5'-TCCCAAACCCTTATGTGAACC (SEQ ID NO:186), the reverse primer is StacR5 5'-GTTTGCCACTCAGAGGAATACTGAAA (SEQ ID NO:187), and the probe is StacP2 5'-CTGCGCCCGGATCCAGGC (SEO ID NO:183).

Standard procedures for the operation of the model 7700, as described in the instrument's manual, were followed using all of the default program settings with the exception of reaction volume which was changed from 50 to 25 ul. Thermal cycling conditions consisted of 2 minutes at 50°C, 10 minutes at 95°C, followed by 40 cycles of 15 seconds at 95°C and 1 minute at 60°C. Cycle threshold ($C_{\rm f}$) determinations were automatically performed by the instrument for each assay using default parameters. Assays for S. chartarum sequences and G. candidum sequences in the same DNA samples were performed in separate reaction tubes.

To quantify conidia the mean S. chartarum calibrator C_T value was subtracted from the mean S. chartarum sequence C_T values in the sample extracts to obtain $\Delta C_{T,STAC}$ values. Ratios of target sequences in the test and calibrator samples were multiplied by the known quantities of S. chartarum conidia in the calibrator samples to obtain measurements of the quantities of these conidia in the test samples. Similar calculations were performed in parallel using G. candidum sequence C_T values from the same calibrator and test samples to

determine $\Delta C_{T,\,GED}$ values and quantities of these conidia in the test samples.

Then G. candidum sequence C_T values were subtracted from mean S. chaztarum sequence C_T values for both test and calibrator sample extracts to obtain ΔC_T values. Calibrator sample ΔC_T values were then subtracted from the test sample ΔC_T values to obtain $\Delta \Delta C_T$ values. These values were used in place of ΔC_T , size values to determine the ratios of S. chartarum target sequences in the test and calibrator samples and to quantify S. chartarum conidia in the test samples as indicated above.

Variances of ΔC_{T} were estimated from the results of the replicate extracts of each sample by:

$$S_{\Delta C\tau}^{2} = S_{Target}^{2} + S_{Ref}^{2} - 2rS_{Target}S_{Ref} [1],$$

where S_{Target} and S_{Ref} are the standard deviations (SD) of the S. chartarum and G. candidum assay results, respectively, and r is the correlation coefficient between these results.

Variances of AAC, were estimated by
$$S_{\text{Lice}}^1 = S_{\text{effect}}^2 + S_{\text{const}}^2$$
 [2],

where $S_{2CT(0)}$ is given by Equation [1] applied to the calibrator results, and $S_{2CT(8)}$ by Equation [1] applied to the test sample results. Since calibrator and test sample C_7 values were

of the C_{r} for the calibrator. Variances of $\Delta C_{r,orc}$ values were calculated in the same manner. Standard errors of difference were determined from the appropriate standard deviation divided by the square root of the number of replicate observations (extractions), and confidence intervals for the differences were constructed using these standard errors.

With N_0 representing the number of cells in the calibrator sample, the corresponding cell numbers in test samples were estimated by $N_02^{-\Delta y}$ [4], where ΔY was the estimator $\Delta\Delta C_T$, $\Delta C_{T,STAC}$, or $\Delta C_{T,GEC}$. In this paper the term "relative error" refers the range implied by one standard deviation about ΔY , i.e. $N_02^{-\Delta Y \pm S_{\Delta Y}}$, in which $S_{\Delta Y}$ is given by equation [2] or [3]. Confidence intervals were constructed around the estimated cell numbers by $N_0^{2-\Delta Y \pm S_{\Delta Y}/\sqrt{3}}$, where t is the appropriate Student t-value and three replicate extractions were used.

In method evaluation experiments, conidia quantities determined by the $\Delta\Delta C_T$, $\Delta C_{T,STAC}$, or $\Delta C_{T,SEO}$ methods (NT) were compared to "known" quantities of conidia added to the dust samples. The "known" quantities were determined from hemosycometer cell counts of three replicate aliquots (at least 400 total counts) of the conidia stock suspensions used for dilution and sample amendment. The "known" value for ΔC_T (N_B) was calculated from equation [4] based on the nemocycometer counts and calculated, and the differences: d = ΔCT - known value were evaluated via analysis

of variance to test the null hypothesis: d=0. The 95% confidence level range for individual observations of d was constructed, assuming d to be normally distributed, and used to characterize the precision of a single estimate utilizing TaqMan quantification. Note that when antilogs are taken, the confidence interval describes lower and upper limits to the ratio $N_{\rm T}/N_{\rm H}$.

The direct enumeration of Stachybotrys conidia in dust samples was performed by weighing dust samples, suspending them in 0.5% Tween 20 to a concentration of 1 mg/ml and, with constant mixing of the suspensions, aliquots were applied to a hemocytometer chamber. Nine replicate aliquots, or fewer if this was sufficient to enumerate at least 400 conidia, were counted in this manner for each suspension. The volumes of the examined grids were used to calculate conidia numbers per ml of suspension and these values converted to numbers per mg of dust. For comparability with relative error of the TagMan estimates, one standard deviation ranges for direct count estimates were calculated. Conidia were assumed to be randomly distributed within each grid. Under this assumption the corresponding relative error is a range such that the observed count represents an observation one standard daviation above or one standard deviation below a Foisson

dust samples taken from two contaminated homes were obtained by $\Delta\Delta C_1$ analyses of TagMan assay results and compared with the results of presumptive direct microscopic enumeration of these conidia. Mean estimates obtained from the TagMan assays fell within, or very close to the 0.24 to 1.04 range of direct counts that was predicted by the method evaluation experiments (Example 2, Table 1).

Example 2, Table 1. Quantities of S. chartarum conidia in home dust samples determined by LACT TaqMan analysis and direct microscopic enumeration.

Location in Home	Proximity to Fungal Growth	<u>ΔΔCT TaqMan Estimate</u> Conidia/5mg dus: Relative Error ^g		<u>Direct count estimate</u> Conidia/5mg dust Relative Error ^g	
Living Rm	Remote Room ^d	6	4 - 10	667	444 - 1000
Basement	Same Room	11001		2333	1877 - 2901
Basement	Same Room	9200	7100 - 12000	26889	25215 - 28674
Dining Rm	Remote Room	560	420 - 740	1444	1096 - 1904
Basement	Same Room	23800	18300 - 31000	30444	28660 - 32340
Basement	Adjacent Room	300	260 - 340	778	534 - 1133
Easement	Same Room ^e	77200	56700 - 105000	68286	50737 - 55597
HVAC		1.7	0.3 - 11.2	556	357 - 866

a Home 1 located in Cleveland, OH

Analyses of known numbers of Stachybotrysconidia over a range f rom 2 x 10^{1} to 2 X 10^{4} in the presence of 10 mg of composite HVAC system dust were found to provide 95% occurrence results within a range from 25% to 104% of expected values using this approach.

A second type of matrix effect that can affect PCR-based analyses of dust samples is the influence of PCR

b Home 2 located in Cincinnati, CH

c Composite HVAC system dust as described in Methods section of text

d Small amount of fungal growth, no confirmed Stachybotrys e Sample collected directly beneath area of fungal growth

inhibitory compounds. Retention of such compounds through the DNA extraction and purification procedures occurred in only one sample in this study. A simple procedure, involving the dilution and re-analysis of a DNA extract from this sample was used to identify this matrix effect and to obtain a corrected estimate of conidia quantities. This procedure should be generally applicable so long as the concentrations of target sequences in the samples are sufficiently high to still be detectable after the inhibitor's effects are negated by dilution. In practice, however, such follow-up analyses are only likely to be necessary when significant differences are observed in the reference sequence assay results of test and calibrator samples in the initial analyses of the samples. Examiple 3: Evaluation of Stachybotrys chartarumin the House of an Infant with Pulmonary Hemorrhage: Quantitative Assessment Before, During and After Remediation

Air samples (Example 3, Table 1) were taken in a home under remediation for mold damage in two ways; either using a cassette filter (37mm with 0.8 mm filter) or with a 810Campler SKC, Fighty Four, FA, connecting to an AirJun-2 High Flow Sampler pump (Gilian Instrument Co., Clearwater, FL) calibrated at a flow rate of 10 liter per min. These samples were taken for a period between 6 and 90 hrs at 10 liter per

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filter).

Conidial stocks of the target fungus, i.e.

Stachybotrys chartarum, and the reference target, i.e.

Geotrichum candidum, were prepared to act as calibrator and internal standard, respectively.

Genomic DNAs were extracted from 20 µl conidial suspensions using a glass bead milling and glass milk adsorption method. Briefly, this method involved mixing test and reference conidia suspensions (10 µl ea.) with 0.3 g of acid-washed glass beads (G-1277; Sigma, St. Louis, MO) and 10 µl, 100 µl and 300 µl, respectively, of glass milk suspension, lysis buffer and binding buffer from an Elu-Quik ENA purification kit (Schleicher and Schuell, Keene, NH; in sterile 2 ml conical bottom, screw cap tubes (506-636; PGC Scientifics, Gaithersburg, MD). The tubes were shaken in a mini beadbeater (Biospec Products, Bartlesville, GE) for one minute at maximum rate and DNAs were recovered in final volumes of 200 µl distilled water after performing a slight modification of the small-scale protocol provided with the Elu-Quik purification kit.

The TagMan probes and primers were obtained from the custom oligonucleotide synthesis facility at PE-Applied Biosystems (Foster City, CA). TagMan probes contained a TAMRA group conjugated to their 3'-terminal nucleotide and a FAM group trinsults their between huckestides as the quencher and reporter fluorochromes, respectively. For Geotrichum

candidum, the forward primer is NS92F: 5'-CACCGCCGTCGCTAC (SEQ ID NO:86), the reverse primer is GcandR1: 5'-AGAAAAGTTGCCCTCTCCAGTT (SEQ ID NO:87), and the probe is GeoP2: 5'-TCAATCCGGAAGCCTCACTAAGCCATT (SEQ ID NO:88). For Stachybotrys chartarum, the forward primer is StacF4 5'-TCCCAAACCCTTATGTGAACC (SEQ ID NO:186), the reverse primer is StacR5 5'-GTTTGCCACTCAGAGAATACTGAAA (SEQ ID NO:187), and the probe is StacP2 5'-CTGCGCCCGGATCCAGGC (SEQ ID NC:188).

pCR reactions were prepared in 0.5 ml thin-walled, optical grade PCR tubes (PE Applied Biosystems, Foster City CA) by addition of the following components: 12.5 µl of TaqMan Universal Master Mix, a 2 x concentrated, proprietary mixture of AmpliTaq Gold DNA polymerase, AmpErase® UNG, dNTPs with UTP, passive reference dye and optimized buffer components (PE Applied Biosystems, Foster City CA); 2.5 µl of a mixture of forward and reverse primers (10 nM each); 2.5 µl of 400 nM TaqMan probe; 2.5 µl of 2 mg/ml bovine serum albumin (fraction V, Sigma Chemical, St. Louis, MG) and 5 µl of DNA template. Standard procedures for the operation of the model 7700, as described in the instrument's manual, were followed. This included the use of all default program settings with the exception of reaction volume which was changed from 50 to 25 µl. Thermal cycling conditions consisting of two min at 50°C,

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reporter dye fluorescence resulting from the synthesis of PCR products to become significantly higher than background fluorescence levels were automatically performed by the instrument for each reaction using default parameters. Assays for S. chartarum (target) sequences and G. candidum (reference) sequences in the same DNA samples were performed in separate reaction tubes.

Results of air sampling with either filters or BicSamplers indicated that the number of airborne S. chartarum spores in this PH house was low before the remediation began (Example 3, Table 1). The number of S. chartarum spores in the air, when the furnace blower was activated (typical condition for the winter months), increased by a factor of 17-47 in the living room. During demolition, the number of S. chartarum spores in the air increased by four orders of magnitude in the basement, about three orders of magnitude in the dining room and about two orders of magnitude in the upstairs bedroom (Example 3, Table 1). Thus this technology, under actual conditions, can detect the target fungus over four orders of magnitude.

Table 1. Results of air sampling for S. chartarum (S.c.) spores in the mold contaminated home.

Cate	Sample Method	Location (Room = RM)	Sample Time (H)	S.c. Spores (#/m² air)
Pre-remediatio	n			
12/29-30	Filter (Passive) ¹	Living Rm	25.5	0.2
	BioSampler (passive)	Living Rm	25.5	0.3
12/30-31	Filter (active) ²	Living Rm	24	9.3
	BioSampler (active)	Living Rm	24	5.0
12/31	Filter (active)	Dining Rm	90	0.1
	BioSample (active)	Dining Rm	90	1.7
12/31 - 1/4	Filter (active)	Easement	90	0.6
During Remedi	iation ³			
1-19	Filter	Basement	6.6	1.1 × 10 ³
	BioSampler	Basement	6.5	1.6 × 10 ³
	Filter PMP ⁴	Basement	6.5	2.0 × 10 ³
1-20	Filter BioSampler Filter PMP	Dining Rm Dining Rm Dining Rm	6.25 6.25 5.75	$ \begin{array}{c} 1.8 \times 10^{3} \\ 2.7 \times 10^{3} \\ 4.0 \times 10^{3} \end{array} $
1-21	Filter	N. Bedrm	7.75	0.1 x 10 ³
	BioSampler	N. Bedrm	7.75	0.1 x 10 ³

^{1 &}quot;passive" means furnace blower off, furnace sealed and inoperable 2 "active" means furnace blower on, furnace operable.

EXAMPLE 4:

Identification and Quantification of Helicobacter pylori

Culturing of Helicobacter pylori from environmental sources continues to be an obstacle in detecting and enumerating this organism. Selection of primer and probe sequences for the urch gene was performed based on comparative sequence analyses of le strains of H. pylori and other Helicobacter species. For Helicobacter pylorii, the forward primer is HpylF1: 5'-GGGTATTGRAGGGATGTTTCCT (SEQ ID NC:223),

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DNA was extracted from aliquots of ten-fold serial dilutions of *H. pylori* by EluQuick kits from Schleeicher and Schuell, Inc. The cells were lysed, DNA bound to glass beads and washed with alcohol and salt reduction solutions followed by elution from filters with water. One set of extraction tubes, contained only *H. pylori*. A second set also received 10³/Ml *E. Coli*. Portions of some DNA extracts were subjected to agarose gel electrophoresis and GelStar staining. Yields of high molecular weight total DNA (appearing as bands on the 1.5% gels) were estimated by comparisons of their fluorescence signals with those of a series of known mass standards (Gibco/BFL) using a model Sl fluorimager (Molecular Dynamics).

The more commonly identified non-pylori Helicobacter species were tested with H. pylori primers and probe (Example 4, Table 1). Results show that when compared to the negative extraction control all of these species were also negative. All obtained C_T values in the, range of 37 to 39. A 40 C_T is the lowest negative value obtainable. Counts of the hacteria were high. They ranged from 107 to 108 per assay. The H. pylo.ri strain also was initially in this range with a cr value was 15.

Example 4, Table 1

Bacteria	Dilution	Cells/TagMan	C _r Values
Campylobacter jejuni	10°	8.75 x 10°	36.55
	10 ⁻¹	8.75 x 10°	36.86
	10 ⁻²	8.75 x 10°	38.78
Helicobacter felis	10°	6.8×10^{1}	37.55
	10-:	6.8×10^{4}	36.17
	10-2	6.8×10^{3}	38.21
Helicobacter hepoticus	10°	2.3×10^{7}	36.68
	10-1	2.3×10^{6}	37.14
	10-2	2.3×10^{5}	39.74
Helicobacter mustelae	10°	1.9 x 10°	34.66
	10 ⁻¹	1.9 X 10°	36.37
	10 ⁻²	1.9 x 10°	37.65
Helicobacter pylori	100 10-1 10-2 10-3 10-6 10-5 10-6 10-7	$\begin{array}{c} 2.1 \times 10^{7} \\ 2.1 \times 10^{6} \\ 2.1 \times 10^{5} \\ 2.1 \times 10^{4} \\ 2.1 \times 10^{3} \\ 2.1 \times 10^{2} \\ 2.1 \times 10^{1} \\ 2.1 \times 10^{1} \end{array}$	14.93 18.23 21.45 25.24 32.73 34.63 35.24 39.58
Negative Extraction Control	-	÷ -	37.65
Positive Calibrator Control	10-1	9.8 x 10 ⁵	17.3

Samples of serially diluted H. pylori cells spanning a 6 log concentration range were subjected to DNA extraction and TagMan analysis.

Estimated cell quantities in the extracted samples ranged from 20 to 2 x 10^6 based on direct microscopic counts following staining with DAPI. Results from 5 replicate experiments showed a good correlation (r2 = 0.99) between

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were seen for two Helicobacter pyloristrains. It was concluded that the TaqMan quantitative PCR method has the potential to provide accurate quantification of H. pyloricells in environmental samples.

Ten-fold dilutions of a single DNA extract are shown in Figure 4 along with the corresponding regression analysis. This curve is linear all the way to a negative $C_{\rm T}$ of 40. The R-squared is 0.999. Counts that correspond to the initial dilution can be extrapolated for the other dilutions and can be included on the X-axis. Figure 4 shows a linear range from 1.5 x 10 5 to 1.5 genome equivalents.

Example 4, Figure 4. Log (base 10) H. pylori counts per assay are plotted against the cycle threshold values.

CONCLUSIONS

Having now fully described this invention, it will be appreciated by those skilled in the art that the same can be performed within a wide range of equivalent parameters, concentrations, and conditions without departing form the spirit and scope of the invention and without undue experimentation.

While this invention has been described in connection with specific embodiments thereof, it will be understood that it is capable of further modifications. This application is intended to cover any variations, uses, or adaptations of the invention following, in general, the principles of the intention and including such departures from

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the present disclosure as come within known or customary practice within the art to which the invention pertains and as may be applied to the essential features hereinbefore set forth as follows in the scope of the appended claims.

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WHAT IS CLAIMED IS:

- 1. A method of detecting and quantifying fungi and bacteria comprising obtaining a sequence of the fungus to be detected and quantified, extracting the DNA from a sample, subjecting said DNA to polymerase chain reaction and fluorescent probe analysis.
- 2. The method according to claim 1 wherein the fungi and bacteria are selected from the group consisting of Absidia coerulea, Absidia glauca, Absidia corymbifera, Acremonium strictum, Alternaria alternata, Apophysomyces elegans, Saksena vasiformis, Aspergillus flavus, Aspergillus oryzae, Aspergillus fumigatus, Neosartoryta fischeri, Aspergillus niger, Aspergillus foetidus, Aspergillus phoenicus, Aspergillus nomius, Aspergillus ochraceus, Aspergillus ostianus, Aspergillus auricomus, Aspergillus parasiticus, Aspergillus sojae, Aspergillus restrictus, Aspergillus caesillus, Aspergillus conicus, Aspergillus sydowii, Aspergillus tamarii, Aspergillus terrues, Asperguters ustus, Aspergillus verricolar, Aspergillus ustus, Aspergillus versicolar, Aspergillus ustus, Aspergillus versicolar, Chaetomium globosum, Cladosporium cladosporicides, Cladosporium herbarum, Cladosporium sphaerospermum,

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amsteledami, Eurotium chevalieri, Eurotium herbarierum, Eurotium ruprum, Eurotium repens, Geotrichum candidum strain UAMH 7863, Geotrichum candidum, Geotrichum klebahrii, Memnoniella echinata, Mortierella polycehpahal, Mortierella wolfii, Mucor mucedc, Mucor amphibiorum, Mucor circinelloides, Mucor heimalis, Mucor indicus, Mucor mucedo, Mucor racemosus, Mucor famosissimus, Phizopus azygosporous, Phizopus homothalicus, Rhizopus microsporus, Rhizopus oligosporus, Rhizopus oryzae, Myrothecium verrucaria, Myrothecium roridum, Paecilomyces lilacinus, Paecilomyces varicti, Penicillium freii, Penicillim verrucosum, Penicillium hirsutum, Penicillium alberechii, Penicillum aurantiogriseum, Penicillium polonicum, Penicillium viridicatum, Penicillium hirsutum, Penicillium brevizompatcum, Penicillium chrysogenum, Fenicillium griseofulvum, Penicillium glandicola, Penicillium coprophilum, Penicillium crustaeceum, Penicillium egyptiacum, Penicililium crustosum, Penicillium citrinum, Penicillium sartoryi, Penicillium westlingi, Penicillium corylophilum, Penicillium decumbens, Fenicillium echinulatum, Penicillium solitum, Penicillium schlerotigenum, Penicillium italicum, Eupenicillium expansum, Penicillium fellutanum, Penicillium charlesii, Penicillium janthinellum, Penicillium raperi, Penicillium madriti, Penicillium gladioli, Penicillium cwalitum, Peraciliium requeicreii, Penicillium simplicissimum, Penicillium cchrochloron, Penicillium spinulcsum, Penicillium

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glabrum. Penicillum thomii, Fenicillium pupurescens,
Eupenicillium lapidosum, Rhizomucor miehei, Rhizomucor
pusillus, Rhizomucor variabilis, Rhizopus stolonifer,
Scopulariopsis asperula, Scopulariopsis brevicaulis,
Scopulariopsis fusca, Scopulariopsis brumptii, Scopulariopsis
chartarum, Scopulariopsis sphaerospora, Trichoderma
aasperellum, Trichoderma hamatum, Trichoderma viride,
Trichoderma harzianum, Trichoderma longibrachiatum,
Trichoderma citroviride, Trichoderma atroviride, Trichoderma
koningii, Ulocladium atrum, Ulocladium chartarum, Uloclacium
botrytis, Wallemia sebi, Escherichia coli, Helicobacter
pylorii, Penicillium verrusosum, and Stachybotrys chartarum.

- 3. The method according to claim 2 wherein the fungi are selected from the group consisting of Absidia ccerulea/glauca, the Forward Primer is NS92F: 5'-CACCGCCGTCGCTAC (SEQ ID NO:1), the reverse primer is AccerR1: 5'-TCTAGTTTGCCATAGTTCTCTTCCAG (SEQ ID NO:2), and the probe is MucF1: 5'-CCGATTGAATGGTTATAGTGAGCATATGGGATC (SEQ ID NO:3).
- 4 . The method according to claim 2 wherein the fungi are selected from the group consisting of Absidia

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- 5. The method according to claim 2 wherein the fungi are selected from the group consisting of Abremonium strictum, the forward primer is AstrcF1: 5'-CAACCCATTGTGAACTTACCAAAC (SEQ ID NO:7), the reverse primer is AstrcR1: 5'-CGCCCCTCAGAGAAATACGATT (SEQ ID NO:8), and the probe is AstrcP1: 5'-TCAGCGCGCGGTGGCCTC (SEQ ID NO:9).
- 6. The method according to claim 2 wherein the fungi are selected from the group consisting of Alternaria alternata, the forward primer is AaltrF1: 5'-3GCGGGCTGGAACCTC (SEQ ID NO:10), the reverse primer is AltrR1-1: 5'-GCAATTACAAAAGGTTTATGTTTGTCGTA (SEQ ID NO:11), or the reverse primer is AaltrR1-2: 5'-TGCAATTACTAAAGGTTTATGTTTGTCGTA (SEQ ID NO:12), and the probe is AaltrP1: 5'-TTACAGCCTTGCTGAATTATTCACCCTTGTCTTT (SEQ ID NO:13).
- 7. The method according to claim 2 wherein the fungi are selected from the group consisting of Apophysomyces elegans and Saksenea vasiformis, the forward primer is NS92F: 5'-CACCGCCCGTCGCTAC (SEQ ID NO:14), the reverse primer is AelegF1: 5'-GACTCGAATGAGTTCTCGCTTC (SEQ ID NO:15), and the probe is AelegF1: 5'-TGGCCAAGACCAGAATATGGGATTGC (SEQ ID NO:16).
- fungi are selected from the group consisting of Aspergillus

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flavus/cryzae, the forward primer is AflavF1: 5'-CGAGTGTAGGGTTCCTAGCGA (SEQ ID NO:17), the reverse primer is AflavR1: 5'-CCGGCGGCCATGAAT (SEQ ID NO:18), and the probe is AflavP1: 5'-TCCCACCCGTGTTTACTGTACCTTAGTTGCT (SEQ ID NO:19).

- 9. The method according to claim 2 wherein the fungi are selected from the group consisting of Aspergillus fumigatus, Neosartorya fischeri, the forward primer is AfumiF1: 5'-GCCGCCGTTTCGAC (SEQ ID NO:20), the reverse primer is AfumiR1: 5'-CCGTTGTTGAAAGTTTTTAACTGATTAC (SEQ ID NO:21), and the probe is AfumiP1: 5'-CCCGCCGAAGACCCCAACATG (SEQ ID NO:22).
- 10. The method according to claim 2 wherein the fungi are selected from the group consisting of Aspergillus niger/foetidus/phoenicus, the forward primeris AnigrF1: 5'-GCCGGAGACCCCAACAC-3' (SEQ ID NO:23), the reverse primer is AnigrR1: 5'-TGTTGAAAGTTTTAACTGATTGCATT-3' (SEQ ID NO:24), and the probe is AnigrP1: 5'-AATCAACTCAGACTGCACGCTTTCAGACAG (SEQ ID NO:C5).
- fungi are selected from the group consisting of Aspergillus nomius, the forward primer is AflavF1: 5'-

- 12 . The method according to claim 2 wherein the fungi are selected from the group consisting of Aspergillus ochraceus/ostianus/auricomus, the forward primer is AcchrF1: 5'-AACCTCCCACCGTGTATACC-3' (SEQ ID NO:29), the reverse primer is AbchrR1: 5'-CCGGCGAGCGCTGTG-3' (SEQ ID NO:30), and the probe is AbchrP1: 5'-ACCTTGTTGGTCGGCGAGCCC (SEQ ID NO:31).
- fungi are selected from the group consisting of Aspergillus parasiticus/sojae, the forward primer is AflavF1: 5'-CGAGTGTAGGGTTCCTAGCGA-3' (SEQ ID NO:32), the reverse primer is AparaR3: 5'-GCCCGGGGCTGACG-3' (SEQ ID NO:33), and the probe is AflavP1: 5'-TCCCACCCGTGTTTACTGTACCTTAGTTGCT (SEQ ID NO:34).
- 14. The method according to claim 2 wherein the fungi are selected from the group consisting of Aspergillus restrictus/daesillus/conicus, the forward primer is ArestF2: 5'-CGGGCCCGCCTTCAT-3' (SEQ ID NO:35), the reverse primer is ArestR1: 5'-GTTGTTGAAAGTTTTAACGATTTTTCT (SEQ ID NO:36), and the probe is ArestP1: 5'-CCCGCCGGAGACTCCAACATTG (SEQ ID NO:37).
- 15. The method according to claim 2 wherein the fungi are selected from the group consisting of Aspergillus APALVIL, the for all primar is Asjdoil. 8 -0-ACCOSTGAP-3' (SEQ ID NO:38), the reverse primer is versR1: 5'-

CCATTGTTGAAAGTTTTGACTGATTTTA (SEQ ID NO:39), and the probe is versP1: 5'-AGACTGCATCACTCTCAGGCATGAAGTTCAG (SEQ ID NO:40).

- 16. The method according to claim 2 wherein the fungi are selected from the group consisting of Aspergillus tamarii, the forward primer is AflavF1: 5'-CGAGTGTAGGGA (SEQ ID NO:41), the reverse primer is AtamaR1: 5'-CCCGGCGGGCCTTAA (SEQ ID NO:42), and the probe is AflavF1: 5'-TCCCACCCGTGTTTACTGTACCTTAGTTGCT (SEQ ID NO:43).
- 17 . The method according to claim 2 wherein the fungi are selected from the group consisting of Aspergillus terreus, the forward primer is AterrF1: 5'TTACCGAGTGCGGGTCTTTA (SEQ ID NO:44), the reverse primer is AterrF1: 5'-CGGCGGCCAGCAAC (SEQ ID NO:45), and the probe is AterrF1: 5'-AACCTCCCACCGTGACTATTGTACCTTG (SEQ ID NO:46).
- 18. The method according to claim 2 wherein the fungi are selected from the group consisting of Aspergillus ustus, the forward primer is AustsF1: t'-GATCATTAGUGAGTGUAGGIUT (SEQ ID NO:47), the reverse primer is AustsF1: 5'-GCCGAAGCAACGTTGGTC (SEQ ID NO:48), and the probe is AustsF1: 5'-CCCCGGGGGGGGGCTAACC (SEQ ID NO:49).

versidolor, the forward primer is AversF2: 5'-CGGCGGGGGGGGCCCT (SEQ ID NO:50), the reverse primer is versR1: 5'-CCATTGTTGAAAGTTTTGACTGATTTTA (SEQ ID NO:51), and the probe is versP1: 5'-AGACTGCACCACTCTCAGGCATGAAGTTCAG (SEQ ID NO:52).

- 20. The method according to claim 2 wherein the fungi are selected from the group consisting of Chaetomium globosum, the forward primer is CglobF1: 5'-CCGCAGGCCCTGAAAAG (SEQ ID NO:53), the reverse primer is CglobR1: 5'-CGCGGGCGCGACCA (SEQ ID NO:54), and the probe is CglobP1: 5'-AGATGTATGCTACTACGCTCGGTGCGACAG (SEQ ID NO:55).
- 21. The method according to claim 2 wherein the fungi are selected from the group consisting of Cladosporium cladosporicides the Type 1, the forward primer is Cclad1F1: 5'-CATTACAAGTGACCCCGGTCTAAC (SEQ ID NO:56), the reverse primer is CcladR1: 5'-CCCCGGAGGCAACAGAG (SEQ ID NO:57), and the probe is CcladP1: 5'-CCGGGGATGTTCATAACCCTTTGTTGTCC (SEQ ID NO:58); and for Type 2 the forward primer is Cclad2F1: 5'-TACAAGTGACCCCGGCTACG (SEQ ID NO:59), the reverse primer is CcladR1: 5'-CCGGGGATGTTCATAACCCTTTGTTGTCC (SEQ ID NO:61).
- 22. The method according to claim 2 wherein the function are selected from the group consisting on Gladesportum. herbarum, the forward primer is CherbF1: 5'-AAGAACGCCCGGGCTT

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(SEQ ID NO:62), the reverse primer is CherbR1: 5'-CGCAAGAGTTTGAAGTGTCCAC (SEQ ID NO:63), and the probe is CherbP1: 5'-CTGGTTATTCATAACCCTTTGTTGTCCGACTCTG (SEQ ID NO:64).

- 23 . The method according to claim 2 wherein the fungi are selected from the group consisting of Cladosporium sphaerospermum, the forward primer is CsphaF1: 5'-ACCGGCTGGGTCTTTCG (SEQ ID NO:65), the reverse primer is CsphaR1: 5'-GGGGTTGTTTTACGGCGTG (SEQ ID NO:66), and the probe is CsphaP1: 5'-CCCGCGGGCACCCTTTAGCGA (SEQ ID NO:67).
- 24. The method according to claim 2 wherein the fungi are selected from the group consisting of Conidiobolus coronatus/incongruus, the forward primer is NS92F: 5'-CACCGCCCGTCGCTAC (SEQ ID NO:68), the reverse primer is ConiR1: 5'-TGACCAAGTTTGACCAATTTCTCTA (SEQ ID NO:69), and the probe is ConiP1: 5'-ATGGTTTAGTGAGGGCCTCTGGATTTGAAGCTT (SEQ ID NO:70).
- IS . The method according to claim 2 wherein the tungs are solerted from the group consisting of Cunninghamella elegans, the corward primer is NSSLF: E'-CACCGCCCGTCGCTAC (SEQ ID NO:71), the reverse primer is CunR1: 5'-AATCTAGTTTGCCATAGTTCTCCTCA (SEQ ID NO:72), and the probe is

- 26. The method according to claim 2 wherein the fungi are selected from the group consisting of Emericella nidulans/rugulosa/quadrilineata, the forward primer is AversF1: 5'-CAACCTCCCACCCGTGAC (SEQ ID NO:74), the reverse primer is AniduR1: 5'-CATTGTTGAAAGTTTTGACTGATTTGT (SEQ ID NO:75), and the probe is versP1: 5'-AGACTGCATCACTCTCAGGCATGAAGTTCAG (SEQ ID NO:76).
- 27 . The method according to claim 2 wherein the fungi are selected from the group consisting of Eurotium mstelodami.'chevalieri/herbariorum/rubrum,'repens, the forward primer is EamstF1: 5'-GTGGCGGCACCATGTCT (SEQ ID NC:77), the reverse primer is EamstR1: 5'-CTGGTTAAAAAGATTGGTTGCGA (SEQ ID NO:78), and the probe is EamstP1: 5'-CAGCTGGACCTACGGGAGCGGG (SEQ ID NO:79).
- The method according to claim 2 wherein the fungi are selected from the group consisting of Epicoccum nigrum, the forward primer is EnigrF1: 5'TTGTAGACTTCGGTCTGCTACCTCTT (SEQ ID NO:80), the reverse primer is EnigrR1: 5'-TGCAACTGCAAAGGGTTTGAAT (SEQ ID NO:31), and the probe is EnigrP1: 5'-CATGTCTTTTGAGTACCTTCGTTTCCTCGGC (SEQ ID NO:82).
- fungi are selected from the group consisting of Geotrichum

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candidum strain UAMH 7863, the forward primer is GeoFl: 5'-GATATTCTTGTGAATPGCAGAAGTGA (SEQ ID NO:83), the reverse primer is GeoRl: 5'-TTGATTCGAAATTTTAGAAGAGCAAA (SEQ ID NO:84), and the probe is GeoPl: 5'-CAATTCCAAGAGAGAAACAACGCTCAAACAAG (SEQ ID NO:85).

- fungi are selected from the group consisting of Geotrichum candidum, the forward primer is NS92F: 5'-CACCGCCCGTCGCTAC (SEQ ID NO:86), the reverse primer is GcandF1: 5'-AGAAAAGTTGCCCTCTCCAGTT (SEQ ID NO:87), and the probe is GeoP2: 5'-TCAATCCGGAAGCCTCACTAAGCCATT (SEQ ID NO:88).
- 31. The method according to claim 2 wherein the fungi are selected from the group consisting of Geotrichum klebahnii, the forward primer is NS92F: 5'-CACCGCCCGTCGCTAC (SEQ ID NO:39), the reverse primer is GklebF1: 5'-AAAASTCGCCCTCTCCTGC (SEQ ID NO:90), and the probe is GeoP2: 5'-TCAATCCGGAAGCCTCACTAAGCCATT (SEQ ID NO:91).
- fungi are selected from the group consisting of Memnoniella echinata, the forward primer is StacF4 5'-

- fungi are selected from the group consisting of Mortierella polycephala/wolfii, the forward primer is NS92F: 5'-CACCGCCCGTCGCTAC (SEQ ID NO:95), the reverse primer is MortR1: 5'-TGACCAAGTTTGGATAACTTTTCAG (SEQ ID NO:96), and the probe is MortP1: 5'-CTTAGTGAGGCTTTCGGATTGGATCTAGGCA (SEQ ID NO:97).
- 34. The method according to claim 2 wherein the fungi are selected from the group consisting of Mucor mucedo, the forward primer is NS92F: 5'-CACCGCCCGTCGCTAC (SEQ ID NO:98), the reverse primer is MmuceR1: 5'-CTAPATATCTAGTTTGCCATAGTTTTCG (SEQ ID NO:99), and the probe is MucF1: 5'-CCGATTGAATGGTTATAGTGAGCATATGGGATC (SEQ ID NO:100).
- fungi are selected from the group consisting of Mucor amphibiorum/circinelloides/heimalis/indicus/mucedo/racemosus/ramosissimus and Rhizopus azygosporus/homothalicus/microsporus/oligosporus/orytae, the forward primer is NS92F: 5'-CACCGCCGTCGCTAC (SEQ ID NO:101), the reverse primer is MucF1-1: 5'-CCTAGTTTGCCATAGTTCTCAGCAG (SEQ ID NO:102), and the probe is MutP1: 5'-CCGATTGARTGGTTATAGTGAGCATATGGGATC (SEQ ID NC:103).
- fungi are selected from the group consisting of Myrothecium

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Verrucaria/roridum, the forward primer is MyroF1: 5'AGTTTACAAACTCCCAAACCCTTT (SEQ ID NO:104), the reverse primer
is MyroR1: 5'-GTGTCACTCAGAGGAGAAACCA (SEQ ID NO:105), and the
probe is MyroP1: 5'-CGCCTGGTTCCGGGCCC (SEQ ID NO:106).

- 37. The method according to claim 2 wherein the fungi are selected from the group consisting of Paecilomyces lilacinus, the forward primer is lilaF1: 5'-CCCACTGTGAACCTTACCTCAG (SEQ ID NO:107), the reverse primer is PlilaR1: 5'-GCTTGTGCAACTCAGAGAAGAAAT (SEQ ID NO:108), and the probe is PlilaP1: 5'-CCGCCCGCTGGGCGTAATG (SEQ ID NO:109).
- J8 . The method according to claim 2 wherein the fungi are selected from the group consisting of Paecilomyces variotii, the forward primer is PvariF1: 5'-CCCGCCGTGGTTCAC (SEQ IDI NO:110) or the forward primer is PvariF2: 5'-CGAAGACCCCTGGAACG (SEQ ID NO:111), and the reverse primer is PvariR1: 5'-GTTGTTGAAAGTTTTAATTGATTGATTGT (SEQ ID NO:112), and the probe is PvariP1: 5'-CTCAGACGGCAACCTTCCAGGGA (SEQ ID NO:113).
- 39. The method according to claim 2 wherein the fungi are selected from the group consisting of Penicillium

- 40 . The method according to claim 2 wherein the fungi are selected from the group consisting of Penicillium aurantiogriseum/polonicum/viridicatum/freii, the forward primer is PauraF2: 5'-ACCGAGTGAGGGGCCCTT (SEQ ID NO:117), the reverse primer is PauraR6: 5'-CCCGGGGGGCCAGTA (SEQ ID NO:118), and the probe is PenP3: 5'-TCCAACCTCCCACCGGTGTTTATTT (SEQ ID NO:119).
- 41 . The method according to claim 2 wherein the fungi are selected from the group consisting of Penicillium brevicompactum*/alberechii, the forward primer is FbrevF1: 5'-CCTTGTTGCTTCGGCGA (SEQ ID NO:120), the reverse primer is PbrevR2: 5'-TCAGACTACAATCTTCAGACAGAGTTCTAA (SEQ ID NO:121), and the probe is PbrevP1: 5'-CCTGCCTTTTGGCTGCCGGG (SEQ ID NO:122).
- 42. The method according to claim 2 wherein the fungi are selected from the group consisting of Penicillium chrysogenum/griseofulvum/glandicola/coprophilum/expansumand Eupenicillium crustaceum/egyptiacum, the forward primer is PchryF1: 5'-CGGGCCCGCCTTAAC (SEQ ID NO:123), the reverse primer is FchryK1-1: 5'-GRAAGTITIAAATAATTTATTTTCACTCAGAGTA (SEQ ID NO:124) or the reverse primer is PchryF2-1: 5'-

GAAAGTTTTAAATAATTTTCACTCAGACCA (SEQ ID NO:125), and the probe is PenPI: 5'-CGCGCCCGCCGAAGACA (SEQ ID NO:126).

- 43. The method according to claim 2 wherein the fungi are selected from the group consisting of Penicillium citrinum/sartoryi/westlingi, the forward primer is PcitrF1: 5'-CCGTGTTGCCCGAACCTA (SEQ ID NO:127), the reverse primer is PcitrR1: 5'-TTGTTGAAAGTTTTAACTAATTTCGTTATAG (SEQ ID NO:128), and the probe is PcitrP2: 5'-CCCCTGAACGCTGTCTGAAGTTGCA (SEQ ID NO:129).
- 44 . The method according to claim 2 wherein the fungi are selected from the group consisting of Penicillium corylophilum, the forward primer is PcoryF1: 5'-GTCCAACCTCCCACCCA (SEQ ID NO:130), the reverse primer is PcoryR3-1: 5'-GCTCAGACTGCAATCTTCAGACTGT (SEQ ID NO:131), and the probe is PcoryP1: 5'-CTGCCCTCTGGCCCGCG (SEQ ID NO:132).
- 45 . The method according to claim 0 wherein the funyi are selected from the group consisting of Penicillium decumbers, the forward primer is Pdecuk3: 5'-GGCCTCCTTTG (SEQ ID NO:133), the reverse primer is Pdecuk3: 5'-AAAAGATTGAIGTGTTCGGCAG (SEQ ID NO:134), and the

fungi are selected from the group consisting of Penicillium echinulatum/solitum/camembertii/commune/crustosum, the forward primer is PchryF1: 5'-CGGGCCGGCGTCTTAAC (SEQ ID NO:136), the reverse primer is PauraR1-1: 5'-GAAAGTTTTAAATTTTTCACTCAGAGTT (SEQ ID NO:137), and the probe is PenP2: 5'-CGCGCCCGGCGAAGACA (SEQ ID NO:138).

- 47 . The method according to claim 2 wherein the fungi are selected from the group consisting of Penicillium expansum/coprophilum, the forward primer is PauraF2: 5'-ACCGAGTGAGGGCCCTT (SEQ ID NO:139), the reverse primer is PchryR6: 5'-CCCGGCGGCCAGTT (SEQ ID NO:140), and the probe is PenP3: 5'-TCCAACCTCCCACCCGTGTTTATTT (SEQ ID NO:141).
- 48. The method according to claim 2 wherein the fungi are selected from the group consisting of Penicillium fellutanum, charlesii, the forward primer is PfellF1: 5'-AACCTCCCACCCGTGTATACTTA (SEQ ID NO:142), the reverse primer is PfellR1: 5'-CTTATCGCTCAGACTGCAAGGTA (SEQ ID NO:143), and the probe is PfellP1: CGGTTGCCCCCGGGGG (SEQ ID NO:144).
- 49. The method according to claim 2 wherein the fungi are selected from the group consisting of Penicillium janthinellum/raperi, the forward primer is PjantF2: 5'1. (A) 1. (II STITIATIONIACOTA (IE. II A):198., the raverse primer is PjantR2: 5'-TTGARAGITTTRACTGATTTAGCTAATCG (SEQ ID NO:146), and

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the probe is FjantP2: 5'-TGCAATCTTCAGACAGCGTTCAGGG (SEQ ID NO:147).

- 50. The method according to claim 2 wherein the fungi are selected from the group consisting of Penicillium madriti/gladioli, the forward primer is PauraF1: 5'-CGGGCCCGCCTTTAC (SEQ ID NO:148), the reverse primer is PchryR1-1: 5'-GAAAGTTTTAAATAATTTATATTTTCACTCAGAGTA (SEQ ID NO:149) or the reverse primer is PchryR2-1: 5'-GAAAGTTTTAAATAATTTTCACTCAGACCA (SEQ ID NO:150), and the probe is PenP2: 5'-CGCGCCCGCCGAAGACA (SEQ ID NO:151).
- 51 . The method according to claim 2 wherein the fungi are selected from the group consisting of Penicillium oxalicum, the forward primer is PoxalF1: 5'-GGGCCCGCCTCACG (SEQ ID NO:152), the reverse primer is PoxalR1: 5'-GTTGTTGAAAGTTTTAACTGATTTAGTCAAGTA (SEQ ID NO:153), and the probe is PoxalP1: 5'-ACAAGAGTTCGTTTGTGTGTCTTCGGCG (SEQ ID NO:154).
- 51 . The method according to claim 2 wherein the fungi are selected from the group consisting of Penicillium requefortii, the forward primer is PchryFl: 5'-CGGGCCCGCCTTAAC

- 53 . The method according to claim 2 wherein the fungi are selected from the group consisting of Penicillium simplicissimum/ochrochloron, the forward primer is FsimpF1-1: 5'-AACCTCCCACCCGTGTTGATT (SEQ ID NO:158), the reverse primer is PsimpR2-1: 5'-SAGATCCGTTGTTGAAAGTTTTATCTG (SEQ ID NO:159) or the reverse primer is PsimpR3-1: 5'-GAGATCCGTTGTTGAAAGTTTTATCTG (SEQ ID NO:160), and the probe is PsimpP1: 5'-CCGCCTCACGGCCGCC (SEQ ID NO:161).
- 54. The method according to claim 2 wherein the fungi are selected from the group consisting of Penicillium spinulosum/glabrum/thomii/pupurescens and Eupenicillium lapidosum, the forward primer is PspinF1: 5'-GTACCTTGTTGCTTCGGTGC (SEQ ID NO:162), the reverse primer is PspinR1: 5'-CGTTGTTGAAAGTTTTAACTTATTTAGTTTAT (SEQ ID NO:163), and the probe is PspinP1: 5'-TCCGCGCGCGCACCGGAG (SEQ ID NO:164).
- 55. The method according to claim 2 wherein the fungi are selected from the group consisting of Rhizemucor miehei/pusillus/variabilis, the forward primer is NS92F: 5'-CACCGCCGTCGCTAC (SEQ ID NO:165), the reverse primer is PmucR1: 5'-GTAGTTTGCCATAGTTCGGCTA (SEQ ID NO:166), and the probe is PmucP1: 5'-TTGAATGGCTATAGTGAGCATATGGGAGGCT (SEQ ID NO:167).
 - 56 . The method according to claim 2 wherein the

fungi are selected from the group consisting of Rhizopus stolonifer, the forward primer is NS92F: 5'-CACCGCCCGTCGCTAC (SEQ ID NO:168), the reverse primer is RstolR1: 5'-GCTTAGTTTGCCATAGTTCTCTAACAA (SEQ ID NO:169), and the probe is MucP1: 5'-CCGATTGAATGGTTATAGTGAGCATATGGGATC (SEQ ID NO:170).

- 57. The method according to claim 2 wherein the fungi are selected from the group consisting of Scopulariopsis asperula, the forward primer is SCbrvF1: 5'-CCCCTGCGTAGTAGATCCTACAT (SEQ ID NO:171), the reverse primer is SCasprR1: 5'-TCCGAGGTCAAACCATGAGTAA (SEQ ID NO:172) and the probe is ScopP1: 5'-TCGCATCGGGTCCCGGCG (SEQ ID NO:173).
- 58. The method according to claim 2 wherein the fungi are selected from the group consisting of Scopulariopsis brevicaulis/fusca, the forward primer is SCbrvF1: 5'-CCCCTGCGTAGTAGATCCTACAT (SEQ ID NO:174), the reverse primer is SCbrvR1: 5'-TCCGAGGTCAAACCATGAAATA (SEQ ID NO:175), and the probe is ScopP1: 5'-TCGCATCGGGTCCCGGCG (SEQ ID NO:176).
- fungi are selected from the group consisting of Scopulariopsis brumptii, the forward primer is SCbrmF1: 5'-

- fungi are selected from the group consisting of Scopulariopsis chartarum, the forward primer is SCchrF1: 5'CCCCCTGCGTAGTAGTAAAGC (SEQ ID NO:18)), the reverse primer is SCchrR1: 5'-TCCGAGGTCAAACCATCAAG (SEQ ID NO:181), and the probe is ScopP1: 5'-TCGCATCGGGTCCCGGGG (SEQ ID NO:182).
- fungi are selected from the group consisting of Scopulariopsis sphaerospora, the forward primer is SCsphFl: 5'CCCCCTGCGTAGTAGTTTACAA (SEQ ID NO:183), the reverse primer is SCsphRl: 5'-CCGAGGTCAAACCATCAAAAG (SEQ ID NO:184), and the probe is ScopPl: 5'-TCGCATCGGGTCCCGGGG (SEQ ID NO:185).
 - fungi are selected from the group consisting of Stachybotrys chartarum, the forward primer is StacF4 5'TCCCAAACCCTTATGTGAACC (SEQ ID NO:136), the reverse primer is StacR5 5'-GTTTGCCACTCAGAGAATACTGAAA (SEQ ID NO:187), and the probe is StacP2 5'-CTGCGCCCCGGATCCAGGC (SEQ ID NO:188).
 - fungi are selected from the group consisting of Trichoderma asperellum/hamatum, the forward primer is TasprF1: 5'totapa consisting of Trichoderma
 TasprR2-1: 5'-GGACTACAGAAAGAGTTTGGTTGCTT (SEQ ID NO:190), and

the probe is TridP1: 5'-CCAAACTGTTGCCTCGGCGGG (SEQ ID NO:191).

- fungi are selected from the group consisting of Trichoderma asperellum/hamatum/viride*, the forward primer is TasprF1: 5'-CCCAAACCCAATGTGAACGT (SEQ ID NO:192), the reverse primer is TasprR1: 5'-TTTGCTCAGAGCTGTAASAAATACG (SEQ ID NO:193), and the probe is TridP1: 5'-CCAAACTGTTGCCTCGGCGGG (SEQ ID NO:194).
- 65. The method according to claim 2 wherein the fungi are selected from the group consisting of Trichoderma harzianum, the forward primer is TharzF1: 5'-TTGCCTCGGCGGGAT (SEQ ID NO:195), the reverse primer is TharzR1: 5'-ATTTTCGAAACGCCTACGAGA (SEQ ID NO:196), and the probe TharzP1: 5'-CTGCCCCGGGTGCGTCG (SEQ ID NO:197).
- - for the week was been about the same with the contract of the

5'-CCCAAACCCAATGTGAACCA (SEQ ID NO:201), the reverse primer is IviriE1: 5'-TCCGCGAGGGGACTACAG (SEQ ID NC:202), and the probe is TridP1: 5'-CCAAACTGTTGCCTCGGCGGG (SEQ ID NO:203).

- fungi are selected from the group consisting of Ulcoladium atrum/chartarum, the forward primer is UatrmF1: 5'-GCGGGCTGGCATCCTT (SEQ ID NO:204), the reverse primer is UatrmR1: 5'-TTGTCCTATGGTGGCGAA (SEQ ID NO:205), and the probe is UloP1: 5'-TGAATTATTCACCCGTGTCTTTTGCGTACTTCT (SEQ ID NO:206).
- fungi are selected from the group consisting of Ulccladium botrytis, the forward primer is UbotrF1: 5'-CCCCCAGCAGTGCGTT (SEQ ID NO:207), the reverse primer is UbotrR1: 5'-CTGATTGCAATTACAAAAGGTTTATG (SEQ ID NO:208), and the probe is UloP1: 5'-TGAATTATTCACCCGTGTCTTTTGCGTACTTCT (SEQ ID NO:209).
- 70. The method according to claim 2 wherein the fungi are selected from the group consisting of Wallemia sebi, the forward primer is WsebiF1: 5'-GGCTTAGTGAATCCTTCGGAG (SEQ ID NO:210), the reverse primer is WsebiR1: 5'-GTTTACCCAACTTTGCAGTCCA (SEQ ID NO:211), and the probe is WsebiF1: 5'-TGTGCCGTTGCCGGCTCAAATAG (SEQ ID NO:211).

- 71 . The method according to claim 2 wherein the fungi are selected from the group consisting of Universal Fungal Group, for ASSAY 1, the forward primer is 5.8F1: 5'-AACTTTCAACAACGGATCTCTTGG (SEQ ID NO:213), the reverse primer is 5.8F1: 5'-GCGTTCAAAGACTCGATGATTCAC (SEQ ID NO:214), and the probe is 5.8P1: 5'-CATCGATGAAGAACGCAGCGAAATGC (SEQ ID NO:215), for ASSAY 2, the forward primer is NS92F: 5'-CACCGCCCGTCGCTAC (SEQ ID NO:216), the reverse primer is ZygR1: 5'-TAATGATCCTTCCGCAGGTTC (SEQ ID NO:217), and the probe is ZygP1: 5'-CCTACGGCAGAACCTTGTTACGACTTTTACTTCCTCTAAA (SEQ ID NO:218).
- 72. The method according to claim 2 wherein the bacteria are selected from the group consisting of Escherichia coli, the forward primer is uidAF1: 5'-GGGCAGGCCAGCGTATC (SEQ ID NO:219), the reverse primer is uidAR1: 5'-CCCACACTTTGCCGTAATGA (SEQ ID NO:220) or the reverse primer is uidAR2: 5'-CGTACACTTTTCCCGGCAAT (SEQ ID NO:221) and the probe is uidAP1: 5'-TGCTGCGTTTCGATGCGGTCA (SEQ ID NO:222).
- 73. The method according to claim 2 wherein the Lagritia are selected from the group consisting of Heliophacter pylorii, the forward primer is HpylF1: 5'-GGGTATTGAAGCGATGTTTCCT (SEQ ID NO:223), the reverse primer is

- 74. The method according to claim 1 wherein the label is a fluorescent label.
- 75. The method according to claim 1 wherein fungi are detected and quantitated using PCE, hybridization, or other molecular techniques.
- 76. The method according to claim 2 wherein the primer and probes are used of determining the cell quantities of fungi and bacteria.
- 77. The method according to claim 2 wherein the fungi are selected from the group consisting of Aspergillus caespitosus the forward primeris AcaesF1:
- 5'-CTCCCACCGTGAATACGTT the reverse primeris AcaesR1:
- 5'-GGCTCAGACGCAACTCTACAAT and the probe is AcaesP1:
- 5'-CACTGTTGCTTCGGCGAGGAGCC.
- 78. The method according to claim 2 wherein the fungi are selected from the group consisting of Aspergillus candidus, the forward primer is AcandF1: 5'-TTACCGAGTGAGGGTTTCTCTGA the reverse primer is Acand R1: 5'-ACAGTGTTCGTGTTGGGGTCTT and the probe is PsimpP1: 5'-CCGCCTCACGGCCGCC.
- 79. The method according to claim 2 wherein the fungi are selected from the group consisting of Aspergillus cervinus,

the forward primer is AdervF1: 5'-CCACCCGTGCTATTGTACCTTT
the reverse primer is AdervF1-1: 5'-CAACTCAGACTGCAATTCAGAACtGT
and the probe is AfumiP2: 5'-TTCTCGGCGGGCGGG.

- 80. The method according to claim 2 wherein the fungi are selected from the group consisting of Aspergillus clavatus, the forward primer is AclavF1: 5'-CCCGCCGTCTTCGGA the reverse primer is AclavF1: 5'-CCGTTGTTGAAAGTTTTAACTGATTATG, and the probe is AfumiP1: 5'-CCCGCCGAAGACCCCAACATG.
- 81. The method according to claim 2 wherein the fungi are selected from the group consisting of Aspergillus flavipes, the forward primer is AflvpF1: 5'-CCACCCGTGACTACTGTACCAC, the reverse primer is AflvpF1: 5'-CCGGCGGCCAGCTAG, the reverse primer is AflvpF2: 5'-AGGCTTTCAGAAACAGTGTTCG, and the probe is AspP1: 5'-TTGCTTCGGCGGGCCCC.
- 82. The method according to claim 2 wherein the fungi are selected from the group consisting of Aspergillus niveus, the forward primer is, AniveF1: 5'-ACCCGTGCCTATTGTACCCT, the reverse primer is AniveR1: 5'-TGCAAACAATCACACTCAGACAC, and the probe is AspF1: 5'-TTGCTTCGGCGGGGCCC.

reverse primer is AochrR2-1: 5'-CGGCGAGCGCTGTtCC, and the probe is AochrP1: 5'-ACCTTGTTGCTTCGGCGAGCCC.

- 84. The method according to claim 2 wherein the fungi are selected from the group consisting of Aspergillus ostianus, the forward primer is AbchrF1: 5'-AACCTCCCACCCGTGTATACC, the reverse primer is AostiR1-1: E'-CGGCGAGCGCTGTTCT, and the probe is AbchrP1: 5'-ACCTTGTTGCTTCGGCGAGCCC.
- E5. The method according to claim 2 wherein the fungi are selected from the group consisting of Aspergillus paradoxus, the forward primer is ApardF1: 5'-CGGGGGGGGTTTACGCT, the reverse primer is ApardR1-1: 5'-GACTGCAACTTCATACAGAGTTGGT, and the probe is PenP2: 5'-CGCGCCCGGAAGACA.
- 86. The method according to claim 2 wherein the fungi are selected from the group consisting of Aspergillus penicillioides, the forward primer is ApeniF2:
 5'-CGCCGGAGACCTCAACC, the reverse primer is ApeniR2:
 5'-TCCGTTGTTGAAAGTTTTAACGA: and the probe is
 ApeniF2: 5'-TGAACACTGTCTGAAGGTTGCAGTCTGAGTATG.
- 87. The method according to claim 2 wherein the fungi are selected from the group consisting of Aspergillus sclerotiorum, the forward primer is AcircF1: 5'-ATTACTGAGTGAGGGTCCCTCG, the reverse primer is AsclrR1:

- 5'-CCTAGGGAGGGGGTTTGA, and the probe is AcircP1:
- E'-CCCGCCGAAGCAACAAGGTACG.
- 88. The method according to claim 2 wherein the fungi are selected from the group consisting of Aspergillus sydowii, the forward primer is AsydoF1-1: 5'-CAACCTCCCACCCGaGAA, the reverse primer is versR1: 5'-CCATTGTTGAAAGTTTTGACTGATTTTA, and the probe is versP1: 5'-AGACTGCATCACTCTCAGGCATGAAGTTCAG.
- F9. The method according to claim 2 wherein the fungi are selected from the group consisting of Aspergillus unguis, the forward primer is AunguF1: 5'-CAACCTCCCACCCTTGAATACT, the reverse primer is AunguR1: 5'-TCACTCTCAGGCATGAAGTTCAG, and the probe is AcaesP1: 5'-CACTGTTGCTTCGGCGAGGAGCC.
- 90. The method according to claim 2 wherein the fungi are selected from the group consisting of Aspergillus wentii, the forward primer is AwentF1:

 5'-CATTACOGRGTGAGGACCTAACC, the reverse primer is AauriR1:

 5'-CGGGGGGCCACGAAT, and the probe is AciroF1:

 5'-CCGGCGGAAGCAACAAGGTACG.
 - 91. The method according to claim 2 wherein the fungi

of the Formacian and Tile to the other than in the graph round of Life is

- 5'-GTCAGAGGCTATAACACACAGCAG, and the probe is CalbP1: 5'-TTTACCGGGCCAGCATCGGTTT.
- 92. The method according to claim 2 wherein the fungi are selected from the group consisting of Candida dubliniensis, the forward primer is CdubF1: 5'-AGATCAGACTTGGTATTTTGCAAGTTA, the reverse primer is CdubR1: 5'-TAGGCTGGCAGTATCGTCAGA, and the probe is CdubP1: 5'-TTTACCGGGCCAGCATCGGTTT.
- 93. The method according to claim 2 wherein the fungi are selected from the group consisting of Candida (Pichia) guilliermondii, the forward primer is CquiF1:

 5'-CCTTCGTGGCGGGGTG, the reverse primer is CquiR1:

 5'-GCAGGCAGCATCAACGC, and the probe is CquiP1:

 5'-CCGCAGCTTATCGGGCCAGC.
- 94. The method according to claim 2 wherein the fungi are selected from the group consisting of Candida haemulonii, the forward primer is ChaeF1: 5'-GGAGCGACAACGAGCAGTC, the reverse primer is ChaeF1: 5'-AGGAGCCAGAAAGCAAGACG, and the probe is ChaeP1: 5'-ATGTAGTACAGCCCTCTGGGCTGTGCA.
- 95. The method according to claim 2 wherein the fungions are relected from the group consisting of Canadia haemplonic type II, the forward primer is Cha2F1: 5'-ATCGGGTGGAGCGGAACT, the reverse primer is Cha2R1: 5'-CGAAGCAGGAACCATCTGAGA, and

the probe is Cha2P1: 5'-AAGTGGGAGCTGATGTAGCAACCCCC.

- 96. The method according to claim 2 wherein the fungi are selected from the group consisting of Candida krusei, the forward primer is CkruF1: 5'-CTCAGATTTGAAATCGTGCTTTG, the reverse primer is CkruF1: 5'-GGGGCTCTCACCCTCCTG, and the probe is CkruF1: 5'-CACGAGTTGTAGATTGCAGGTTGGAGTCTG.
- 97. The method according to claim 2 wherein the fungi are selected from the group consisting of Candida lipolytica, the forward primer is ClipF1: 5'-TAGCGAGACGAGGGTTACAAATG, the reverse primer is ClipF1: 5'-CGTCGGTGGCAGTGTGGA, and the probe is ClipP1: 5'-CCTTCGGGCGTTCTCCCCTAACC.
- 98. The method according to claim 2 wherein the fungi are selected from the group consisting of Candida lusitaniae, the forward primer is ClusF1: 5'-GGGCCAGCGTCAAATAAAC, the reverse primer is ClusR1: 5'-CGCAGGCCTCAAACAACA, and the probe is ClusF1: 5'-AGAATGTGGCGTGCCTTCG.
- 99. The method according to claim 2 wherein the fungi are selected from the group consisting of Candida maltosa, the forward primer is CmalF1: 5'-GGCCAGCATCAGTTTGGAC,

- 101. The method according to claim 2 wherein the fungi are selected from the group consisting of Candida parapsilosis, the forward pPrimer is CparF1: 5'-GATCAGACTTGGTATTTTGTATGTTACTCTC, the reverse primer is CparR1: 5'-CAGAGCCACATTTCTTTGCAC, and the probe is CparP1: 5'-CCTCTACAGTTTACCGGGCCAGCATCA.
- 101. The method according to claim 2 wherein the fungi are selected from the group consisting of Candida sojae, the forward primer is CscjFl: 5'-CGGTTGTGTTATAGCCTTCGTA, the reverse primer is CscjRl: 5'-ATCATTATGCCAACATCCTAGGTAAT, and the probe is CtropP2: 5'-CGCAGTCCTCAGTCTAGGCTGGCAG.
- 102. The method according to claim 2 wherein the fungi are selected from the group consisting of *Candida tropicalis*, the forward primer is CtropF1:
 5'-GCGGTAGSAGAATTGCGTT, the reverse primer is CtropR2:
 5'-TCATTATGCCAACATCCTAGGTTTA, and the probe is
 CtropF2: 5'-CGCAGTCCTCAGTCTAGGCTGGCAG.
- 103. The method according to claim 2 wherein the fungi are selected from the group consisting of Candida wiswarsthis, the forward primes is CwisFi: 5'-0.880A03A03ATCGGGT the reverse primer is CwisRi: 5'-TCTAGGCTGGCAGTATCCACG, and the probe is CwisFi: 5'-AATGTGGCACGGCCTCGGC.

- 104. The method according to claim 2 wherein the fungi are selected from the group consisting of Candida zeylanoides, the forward primer is Czey F1:
 5'-GTTGTAATTTGAAGAAGGTAACTTTGATT, the reverse primer is Czey R1:
 5'-GACTCTTCGAAAGCACTTTACATGG, and the probe is Czey P1:
 5'-CCTTGGAACAGGACGTCACAGAGGGT.
- 105. The method according to claim 2 wherein the fungi are selected from the group consisting of Emericella (Aspergillus) nidulans/rugulosa/quadrilineata, the forward primer is AversF1: 5'-CAACCTCCCACCCGTGAC, the reverse primer is AniduR1-1: 5'-CCATTGTTGAAAGTTTTGACTGATaTGT, and the probe is versP1: 5'-AGACTGCATCACTCTCAGGCATGAAGTTCAG.
- 106. The method according to claim 2 wherein the fungi are selected from the group consisting of *Geotrichum klebahnii*, the forward primer is GklebF1: 5'-GGGCGACTTTTCCGGC, the reverse primer is GklebR2: 5'-TGGCACREATTCTCCTCTARTTTATTTA, and the prope is GklebF1:
 - 107. The method according to claim 2 wherein the

5'-AAGCTAGTCAAACTTGGTCATTTAGAGGAAGTAAAAGTC.

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- 5'-GAAAGTTTTAAATAATTTATTTTCACTCAGAGTA, and the probe is PenP2: 5'-CGCGCCCGCCGAAGACA.
- 108. The method according to claim 2 wherein the fungi are selected from the group consisting of *Penicillium atramentosum*, the forward primer is PgrisF1-1: 5'-ACCTGCGGAAGGATCATTCT, the reverse primer is PatraR1: 5'-CCCCGGCGGCCCATA, the probe is PenP3: 5'-TCCAACCTCCCACCCGTGTTTATTT.
- 109. The method according to claim 2 wherein the fungi are selected from the group consisting of *Penicillium aurantiogriseum*, the forward primer is PauraF3:
 5'-CGCCGGGGGGCTTC, the reverse primer is PauraR1-1:
 5'-GAAAGTTTTAAATAATTTATATTTTCACTCAGAGTT, the probe is PenP2: 5'-CGCGCCCGCCGAAGACA.
- 110. The method according to claim 2 wherein the fungi are selected from the group consisting of Penicillium aurantiogriseum/polonicum/viridicatum/freii, the forward primer is PexpaFl-1: 5'-TTACCGAGTGAGGGCCGTT, the reverse primer is PauraR6: 5'-CCCGGCGGCCAGTA, the probe is PenP3: 5'-TCCAACCTCCCACCCGTGTTTATTT.
- 111. The method according to claim 2 wherein the fungi are selected from the group consisting of Fenicillium

canescens, the forward primer is PcaneF1:

- 5'-TTACCGAGCGAGAATTCTCTGA, the reverse primer is PcaneR1:
- 5'-AGACTGCAATTITCATACAGAGTTCA, the probe is

PsimpP1: 5'-CCGCCTCACGGCCGCC.

- 112. The method according to claim 2 wherein the fungi are selected from the group consisting of *Fenicillium citreonigrum*, the forward primer is PcteoF1-1:

 5'-TGTTGGGCTCCGTCCTCtTC, he reverse primer is PcteoR1-1:
- 5'-CGGCCGGGCCTtCAG, the probe is PenP7: 5'-CCGAAAGGCAGCGGCGGC.
- 113. The method according to claim 2 wherein the fungi are selected from the group consisting of *Penicillium coprophilum* the forward primer is PcoprFl-1:
- 5'-GGGTCCAACCTCCCACtCA, the reverse primer is PchryR1-1:
- 5'-GAAAGTTTTAAATAATTTATATTTTCACTCAGAgTA, the probe is PenP1:
- 5'-CGCCTTAACTGGCCGCCGG.
- 114. The method according to claim 2 wherein the fungi are selected from the group consisting of *Ponicillium* crustosum, the forward primer is PorusF1: 5'-CGCCGGGGGGCTTA, the reverse primer is PauraR1-1:
- 5'-GRAAGTTTTARATAATTTATATTTTCACTCAGAGTT,

- 115. The method according to claim 2 wherein the fungi are selected from the group consisting of *Penicillium digitatum*, the forward primer is PaethF1-1: 5'-CGGGGGGGCTCtCGCT, the reverse primer is PdigiR1: 5'-CGTTGTTGAAAGTTTTAAATAATTTCGT, the probe is PenP2: 5'-CGCGCCCGCGAAGACA.
- 116. The method according to claim 2 wherein the fungi are selected from the group consisting of *Penicillium* expansum, the forward primer is PempaF2-1:
- 5'-TCCCACCGTGTTTATTTAGaTC, the reverse primer is PexpaR1:
- 5'-TCACTCAGACGACAATCTTCAGG or PempaR1-1:
- 5'-TCACTCAGACGACAATCTTCtGG, the probe is PenP1:
- 5'-CGCCTTAACTGGCCGCCGG.
- 117. The method according to claim 2 wherein the fungi are selected from the group consisting of *Penicillium freeii*, the forward primer is PfreiF1: 5'-TCACGCCCCCGGGT, the reverse primer is PauraR1-1:
- 5'-GAAAGTTTTAAATTAATTTATATTTCACTCAGAGTT, the probe is PenP2: 5'-CGCGCCCGCCGAAGACA.
- 118. The method according to claim 2 wherein the fungi are selected from the group consisting of Penicillium glandicula, the forward primer is FglanFI-1.
- 5'-CCGGGGGGCTTtCGT, the reverse primer is PchryR1:

- 5'-GAAAGTTTTAAATAATTTATATTTTCACTCAGACTA, the probe is FenP2: 5'-CGCGCCCGCCGAAGACA.
- 119. The method according to claim 2 wherein the fungi are selected from the group consisting of *Penicillium griseofulvum*, the forward primer is PgrisF1-1:
- 5'-ACCTGCGGAAGGATCATTtCT, the reverse primer is PchryR6:
- 5'-CCCGGCGCCAGTT, the probe is PenP3:
- 5'-TCCAACCTCCCACCCGTGTTTATTT.
- 120. The method according to claim 2 wherein the fungi are selected from the group consisting of *Penicillium hirsutum**, the forward primer is PhirsF1-1: 5'-GCCGGGGGGCTCAtA, the reverse primer is PauraR1-1:
- 5'-GAAAGTTTTAAATAATTTATTTTCACTCAGAgTT, the probe is PenP2: 5'-CGCGCCCGCCGAAGACA.
- 121. The method according to claim 2 wherein the fungi are selected from the group consisting of Penicillium implicatum, the forward primer is PimplF1: 5'-GCCGAAGACCCCCCTGT, the reverse primer is PimplR1: 5'CGTTGTTGAAAGTTTTGACTGATTGT, the probe is PimplP1: 5'-AACGCTGTCTGAAGCTTGCAGTCTGAGC.

the reverse primer is PislaR1: 5'-GGCAACGCGGTAACGGTAG, the probe is PislaP1: 5'-AGCCCAACCTCCCACCGTG.

- 123. The method according to claim 2 wherein the fungi are selected from the group consisting of *Penicillium italicum*, the forward primer is PitalF1-1:
- 5'-CTCCCACCCGTGTTTATTTAtCA, the reverse primer is PexpaR1:
- 5'-TCACTCAGACGACAATCTTCAGG or FexpaR1-1:
- 5'-TCACTCAGACGACAATCTTCtGG, the probe is PenP1: (+)
- 5'-CGCCTTAACTGGCCGCCGG.
- 124. The method according to claim 2 wherein the fungi are selected from the group consisting of *Penicillium melinii*, the forward primer is PmeliF1-1:
- 5'-CACGGCTTGTGTGTTGGtCT, the reverse primer is PmeliR1: .
- 5'-GGGCCTACAAGAGCGGAA, the probe is PenP7: 5'-CCGAAAGGCAGCGGCGGC.
- 125. The method according to claim ? wherein the fungi are selected from the group consisting of *Penicillium miczynskii*, the forward primer is PmiczFl-1:
- 5'-GTGTTTAACGAACCTTGTTGCaTT, the reverse primer is PmiczR1-1:
- 5'-CTCAGACTGCATACTTCAGACaGA, the probe is PsimpP1:
- 5'-CCGCCTCACGGCCGCC.

- 126. The method according to claim 2 wherein the fungi are selected from the group consisting of *Penicillium olsonii*, the forward primer is PolsnFl: 5'-GGCGAGCCTGCCTTCG, the reverse primer is PenR2:
- 5'-GATCCGTTGTTGAAAGTTTTAAATAATTTATA, the probe is PolsnP2: 5'-TCCGCGCTCGCCGGAGAC.
- 127. The method according to claim 2 wherein the fungi are selected from the group consisting of Penicillium purpurogenum, the forward primer is PpurpF1:
- 5'-AGGATCATTACTGAGTGCGGA, the reverse primer is PpurpR1: 5'-GCCAAAGCAACAGGGTATTC, the probe is PpurpP1: 5'-CCCTCGCGGGGTCCAACCTCC.
- 128. The method according to claim 2 wherein the fungi are selected from the group consisting of *Penicillium* raistrickii, the forward primer is PgrisF1-1:
- 5'-ACCTGCGGAAGGATCATTtCT, the reverse primer is PraisR1:
- 5'-CCCGGCGGCCAGAC, the probe is PenP3:
- E'-TucaAboloccarerTTATTI.
- 129. The method according to claim2 wherein the fungi are selected from the group consising of Fenicillium restrictum,

- 130. The method according to claim 2 wherein the fungi are selected from the group consisting of *Penicillium sclerotiorum*, the forward primer is P sclrF1:
 5'-TTCCDCCGGGGAACAGG, the reverse primer is P sclrR1:
 5'-GCCCCATACGCTCGAGGAT, the probe is P sclrP1: 5'CCGAAAGGCAGTGGCGGCAC.
- 131. The method according to claim 2 wherein the fungi are selected from the group consisting of *Penicillium* simplicissimum/ochrochloron, the forward primer is PsimpF2-1: 5'-CGCCGAAGACACCATTGAtCT, the reverse primer is PsimpR4-1: 5'-CTGAATTCTGCAATTCACATAACG, the probe is PsimpP2: 5'-TGTCTGAAGATTGCAGTCTGAGCGATTAGC.
- 132. The method according to claim 2 wherein the fungi are selected from the group consisting of *Penicillium variabile*, the forward primer is PvarbF1: 5'-GCCGGGGGGCTTCT, the reverse primer is PvarbR1: 5'-TCTCACTCAGACTCACTGTTCAGG, the probe is PvarbP1: 5'-AGGSTTCTAGGGTGCTTCGGCGG.
- 133. The method according to claim 2 wherein the fungi are selected from the group consisting of *Penicillium varrucosum**, the forward primer is PverrF2: 5'-CGGGCCCGCCTTTG, the newscape number is FauraR1'.
- 5'-GAAAGTTTTAAATATTTTCACTCAGACTT, the probe is PenP2: 5'-CGCGCCCCGCAAGACA.

- 134. The method according to claim 2 wherein the fungi are scleeted from the group consisting of *Penicillium* waksmanii, the forward primer is P waksF1-1:
 5'-GTGTTTAACGAACCTTGTTGCATC, the reverse primer is P waksR1-1:
 5'-CTTCAGACAGCGTTCACAGGTAG, the probe is PsimpP1:
 5'-CCGCCTCACGGCCGCC.
- 135. The method according to claim 2 wherein the fungi are selected from the group consisting of *Ulocladium* atrum, the forward primer is UatrmF2: 5'-CGGGCTGGCATCCTTC, the reverse primer is UatrmR2: 5'-CTGATTGCAATTACAAAAGGTTTATG, the probe is UloP1: 5'-TGAATTATTCACCCGTGTCTTTTGCGTACTTCT.
- 136. The method according to claim 2 wherein the fungi are selected from the group consisting of *Ulocladium chartarum*, the forward primer is UcharF1-1:
 5'-AGCGGGCTGGAATCCaTT, the reverse primer is UcharR1-1:
 5'-CTGATTGCAATTACAAAAGGTTGAAT, the probe is UlcP1:
 5'-TGAATTATTCACCCGTGTCTTTTGCGTACTTCT.
- 137. The method according to claim 2 wherein the bacteria are selected from the group consisting of Legionella maceachernii, the forward primer is LmaceF1:

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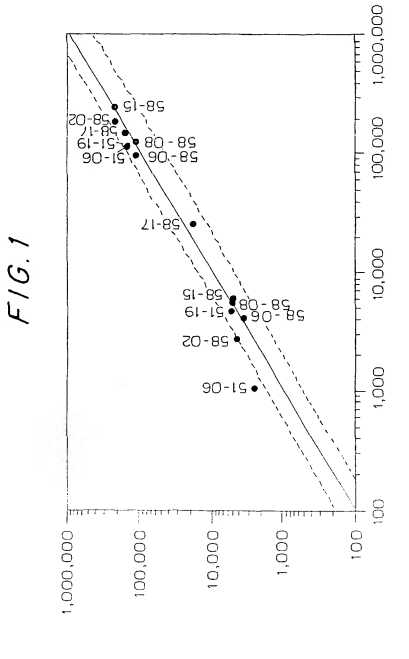
138. The method according to claim 2 wherein the bacteria are selected from the group consisting of Legionella micdadei, the forward primer is LmicdF1:

5'-GGTGGTTTTATAAGTTATCTGTGAAATTC the reverse primer is

PmiodR1: 5'-CACTACCCTCTCCTATACTCAAAGTCTC the probe is

LmicdPl: 5'-AGTCTTATCTGACCACCCTAGGTTGAGCCCA.

- 139. The method according to claim 2 wherein the bacteria are selected from the group consisting of Legionella pneumophila the forward primer is LpneuF1:
- 5'-CGGAATTACTGGGCGTAAAGG the reverse primer is PpneuR1:
- 5'-GAGTCAACCAGTATTATCTGACCGT the probe is LpneuP1: 5'-
- 140. The method according to claim 2 wherein the bacteria are selected from the group consisting of Legionella sainthelensi/cincinnatiensis, the forward primer is LsainF1: 5'-CGTAGGAATATGCCTTGAAGACT the reverse primer is PsainR1: 5'-AAGGTCCCCAGCTTTCGT the probe is LsainP1: 5'-AGACATCATCCGSTATTAGCTTGAGTTTCCC.

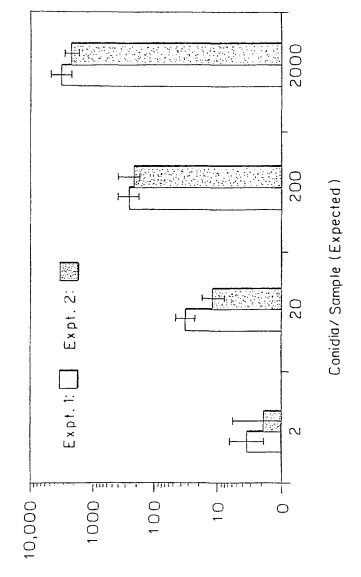


Conidia / Sample (Hemocytometer Count)

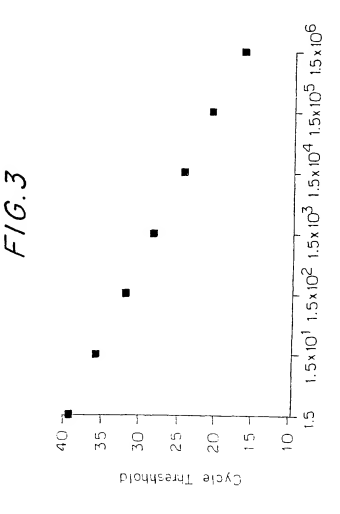
Line of Equality
(TaqMan = Hemocytometer)

95% Confidence Range for Observations F16.2

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Conidia / Sample (TaqMan Estimate)



H.pylori DNA / Assay (genome equivalents)

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